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(54) Title: MODULATORS OF TISSUE REGENERATION

(57) Abstract

Proteins which are upregulated in injured or regenerating tissues, as well as the DNA encoding these proteins, are disclosed, as well as therapeutic compositions and methods of treatment encompassing these compounds.

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MODULATORS OF TISSUE REGENERATION

FIELD OF THE INVENTION

The invention relates to proteins which are upregulated in injured or regenerating tissues, as well as to the DNA encoding these proteins. The invention further relates to the repeutic compositions and methods of treatment encompassing these proteins.

5 **BACKGROUND OF THE INVENTION**

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A dynamic remodeling of tissue architecture occurs during development and during tissue repair after injury. To study this process, we have focused on a model of kidney injury caused by an ischemia-reperfusion insult.

The kidney is able to repair damage to the proximal tubule epithelium through a complex series of events involving cell death, proliferation of surviving proximal tubule epithelial cells, formation of poorly differentiated regenerative epithelium over the denuded basement membrane, and differentiation of the regenerative epithelium to form a fully functional proximal tubule epithelial cells (Wallin et al., Lab. Invest. 66:474-484, 1992; Witzgall et al., Mol. Cell. Biol. 13:1933-1942, 1994; Ichimura et al., Am. J. Physiol. 269:F653-662, 1995; Thadhani et al., N. Engl. J. Med. 334:1448-1460, 1996). Growth factors such as IGF, EGF, and HGF have been implicated in this process of repair, as has the endothelial cell adhesion molecule ICAM-1. However, the mechanisms by which the tubular epithelial cells are restored are still not understood.

To identify molecules involved in process of injury and repair of the tubular epithelium, we analyzed the difference in the mRNA populations between injured/regenerating and normal kidneys using representational difference analysis (RDA). RDA is a PCR-based method for subtraction which yields target tissue or cell specific cDNA fragments by repetitive subtraction and amplification (Hubank and Schutz, Nucl. Acids Res. 22:5640-5648, 1994).

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SUMMARY OF THE INVENTION

The invention generally provides Kidney Injury-related Molecules (each of which is henceforth called a "KIM") which are upregulated in renal tissue after injury to the kidney. The KIM proteins and peptides of the invention, as well as their agonists and antagonists, and their corresponding are useful in a variety of therapeutic interventions.

The invention provides a purified and isolated DNA molecule having a nucleotide sequence set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6. The invention also includes the complementary strands of these sequences, DNA molecules which hybridize under stringent conditions to the aforementioned DNA molecules, and DNA molecules which, but for the degeneracy of the genetic code, would hybridize to any of the DNA molecules defined above. These DNA molecules may be recombinant, and may be operably linked to an expression control sequence.

The invention further provides a vector comprising a purified and isolated DNA molecule having a nucleotide sequence set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6, or one of the other DNA molecules defined above. This vector may be a biologically functional plasmid or viral DNA vector. One embodiment of the invention provides a prokaryotic or eukaryotic host cell stably transformed or transfected by a vector comprising a DNA molecule of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6. In another embodiment of the invention, a process is provided for the production of a KIM polypeptide product encoded by a DNA molecule as described above; the process involves growing, under suitable culture conditions, prokaryotic or eukaryotic host cells transformed or transfected with the DNA molecule in a manner allowing expression of the DNA molecule, and recovering the polypeptide product of said expression.

A purified and isolated human KIM protein substantially free of other human proteins is specifically within the invention, as is a process for the production of a polypeptide product having part or all of the primary structural conformation and the biological activity of a KIM protein. KIM proteins of the invention may have an amino acid sequence which comprises SEQ ID NO:3, SEQ ID NO:5, or SEQ ID NO:7, or may be a variant of SEQ ID NO:3, SEQ ID NO:5 or SEQ ID NO:7, or a purified and isolated protein encoded by the DNA of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6. These proteins can be provided substantially free of other human proteins. The invention further includes variants of these proteins, such as soluble

variants or fusion proteins. KIM fusion proteins of the invention may comprise an immunoglobulin, a toxin, an imageable compound or a radionuclide.

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The invention also provides a specific monoclonal antibody to the KIM proteins described above. The anti-KIM antibody may be associated with a toxin, imageable compound or radionuclide. Further taught is a hybridoma cell line which produces such a specific antibody.

Pharmaceutical compositions are also within the scope of the invention. A pharmaceutical composition of the invention may comprise a therapeutically effective amount of a KIM protein or anti-KIM antibody of the invention, along with a pharmacologically acceptable carrier.

Diagnostic methods are within the invention, such as assessing the presence or course of resolution of renal injury by measuring the concentration of KIM in urine, serum, or urine sediment of patients who have or who are at risk of developing renal disease.

Methods of treatment of the invention include treating patients with therapeutically effective amounts of KIM, KIM variants, KIM analogs, KIM fusion proteins, KIM agonists, and antibodies to KIM or to KIM ligands. Other therapeutic compounds of the invention include KIM ligands, anti-KIM antibodies, and fusions proteins of KIM ligands. These compounds can be useful in therapeutic methods which either stimulate or inhibit cellular responses that are dependent on KIM function.

Further methods of the invention inhibit the growth of KIM-expressing tumor cells by contacting the cells with a fusion protein of a KIM ligand and either a toxin or radionuclide, or with an anti-KIM antibody conjugated to a toxin or to a radionuclide. Likewise, growth of tumor cells which express KIM ligand may be inhibited by contacting the cells with a fusion protein of a KIM and either a toxin or radionuclide, or with an anti-KIM ligand antibody conjugated to a toxin or to a radionuclide.

The invention also encompasses methods of gene therapy. These include a method of treating a subject with a renal disorder, a method of promoting growth of new tissue in a subject, and a method of promoting survival of damaged tissue in a subject, comprising administering to the subject a vector which includes DNA comprising the nucleotide sequence of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6.

The compounds of the invention are also useful for imaging tissues, either in vitro or in vivo. One such method involves targeting an imageable compound to a cell expressing a protein

-4-

of SEQ ID NO:3, SEQ ID NO:5 or SEQ ID NO:7, comprising contacting the cell with either a monoclonal antibody of the invention or a fusion protein comprising a protein as described above, fused to an imageable compound. For *in vivo* methods, the cell is within a subject, and the protein or the monoclonal antibody is administered to the subject.

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The invention also includes diagnostic methods, such as a method of identifying damage or regeneration of renal cells in a subject, comprising comparing the level of expression of either SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6 in renal cells of the subject to a control level of expression of the sequence in control renal cells. Another method of the invention includes identifying upregulation of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6 in cells comprising contacting the cells with an antisense probe and measuring hybridization to RNA within the cell.

A further embodiment of the diagnostic methods of the invention includes assessing the presence or concentration of a molecule of the invention either in urine, serum, or other body fluids, or in urine sediment or tissue samples. The measured injury-related molecule can be correlated with the presence, extent or course of a pathologic process. This correlation can also be used to assess the efficacy of a therapeutic regime.

BRIEF DESCRIPTION OF THE DRAWINGS

FIGURE 1 is the nucleotide sequence of rat clone cDNA 3-2, with putative protein reading frame of 615 to 1535.

FIGURE 2 is a listing of the cDNA sequence of rat clone 1-7, with putative protein reading frame of 145 to 1065.

FIGURE 3 is a listing of the cDNA sequence of rat clone 4-7, with putative protein reading frame of 107 to 1822.

FIGURE 4 is a listing of the cDNA and deduced amino acid sequences of human clone HI3-10-85, with putative protein reading frame of 1 to 1002. The upper line of the listing is the cDNA

-5-

sequence (SEQ ID NO:6), and the lower line is the deduced amino acid sequence (SEQ ID NO:7).

FIGURE 5 is a BESTFIT comparison of the nucleotide sequence of human clone HI3-10-85 with that of rat clone 3-2.

5 **DETAILED DESCRIPTION OF THE INVENTION**

We identified KIM genes by analyzing differences in mRNA expression between regenerating and normal kidneys using representational difference analysis (RDA). RDA is a PCR-based method for subtraction which yields target tissue or cell-specific cDNA fragments by repetitive subtraction and amplification. The cDNA representation from 48 hr postischemic adult rat kidney RNA is subtracted with the sample from normal (sham-operated) adult rat kidney. In this procedure, sequences which are common to both postischemic and to normal kidney samples are removed, leaving those sequences which are significantly expressed only in the injured kidney tissue. Such genes encode proteins that may be therapeutically beneficial for renal disorders or involved in the injury process. Several clones have been obtained, sequenced and characterized. The clones are then investigated for their expression patterns during kidney repair, development and tissue distribution by northern analysis and RNA in situ hybridization.

Sequence Identification Numbers

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Nucleotide and amino acid sequences referred to in the specification have been given the following sequence identification numbers:

20 SEQ ID NO:1 - nucleotide sequence of rat 3-2 cDNA insert

SEQ ID NO:2 - nucleotide sequence of rat 1-7 cDNA insert

SEQ ID NO:3 - amino acid sequence of rat KIM-1, encoded by rat 3-2 and 1-7 cDNA's

SEQ ID NO:4 - nucleotide sequence of rat 4-7 cDNA insert

SEQ ID NO:5 - amino acid sequence encoded by 4-7 cDNA insert

25 SEQ ID NO:6 - nucleotide sequence of human cDNA clone H13-10-85

SEQ ID NO:7 - amino acid sequence encoded by human cDNA clone H13-10-85

-6-

Definitions of Terms

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A "KIM protein", herein used synonymously with "KIM", is a protein encoded by mRNA which is selectively upregulated following injury to a kidney. One group of KIM proteins of interest includes those coded for by mRNA which is selectively upregulated at any time within one week following any insult which results in injury to renal tissue. Examples of times at which such upregulation might be identified include 10 hours, 24 hours, 48 hours or 96 hours following an insult. Examples of types of insults include those resulting in ischemic, toxic or other types of injury.

A "KIM agonist" is a molecule which can specifically trigger a cellular response normally triggered by the interaction of KIM with a KIM ligand. A KIM agonist can be a KIM variant, or a specific antibody to KIM, or a soluble form of the KIM ligand.

A "KIM antagonist" is a molecule which can specifically associate with a KIM ligand or KIM, thereby blocking or otherwise inhibiting KIM binding to the KIM ligand. The antagonist binding blocks or inhibits cellular responses which would otherwise be triggered by ligation of the KIM ligand with KIM or with a KIM agonist. Examples of KIM antagonists include certain KIM variants, KIM fusion proteins and specific antibodies to a KIM ligand or KIM.

A "KIM ligand" is any molecule which noncovalently and specifically binds to a KIM protein. Such a ligand can be a protein, peptide, steroid, antibody, amino acid derivative, or other type molecule, in any form, including naturally-occurring, recombinantly produced, or otherwise synthetic. A KIM ligand can be in any form, including soluble, membrane-bound, or part of a fusion construct with immunoglobulin, fatty acid, or other moieties. The KIM ligand may be an integrin. A membrane-bound KIM ligand can act as a receptor which, when bound to or associated with KIM, triggers a cellular response. In some interactions, KIM may associate with more than a single KIM ligand, or may associate with a KIM ligand as part of a complex with one or more other molecules or cofactors. In a situation where both the KIM and the KIM ligand are bound to cell membranes, the KIM may associate and react with KIM ligand which is bound to the same cell as the KIM, or it may associate and react with KIM ligand be bound to a second cell. Where the KIM ligation occurs between molecules bound to different cells, the two cells may be the same or different with respect to cellular type or origin, phenotypic or metabolic condition, or type or degree of cellular response (e.g., growth, differentiation or apoptosis) to a given stimulus. "KIM ligation" refers to the contact and binding of KIM with a KIM ligand.

-7-

By "alignment of sequences" is meant the positioning of one sequence, either nucleotide or amino acid, with that of another, to allow a comparison of the sequence of relevant portions of one with that of the other. An example of one method of this procedure is given in Needleman et al. (J. Mol. Biol. 48:443-453, 1970). The method may be implemented conveniently by computer programs such as the Align program (DNAstar, Inc.). As will be understood by those skilled in the art, homologous or functionally equivalent sequences include functionally equivalent arrangements of the cysteine residues within the conserved cysteine skeleton, including amino acid insertions or deletions which alter the linear arrangement of these cysteines, but do not materially impair their relationship in the folded structure of the protein. Therefore, internal gaps and amino acid insertions in the candidate sequence are ignored for purposes of calculating the level of amino acid sequence homology or identity between the candidate and reference sequences. One characteristic frequently used in establishing the homology of proteins is the similarity of the number and location of the cysteine residues between one protein and another.

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"Antisense DNA" refers to the sequence of chromosomal DNA that is transcribed.

An "antisense probe" is a probe which comprises at least a portion of the antisense DNA for a nucleic acid portion of interest.

By "cloning" is meant the use of in vitro recombination techniques to insert a particular gene or other DNA sequence into a vector molecule. In order to successfully clone a desired gene, it is necessary to employ methods for generating DNA fragments, for joining the fragments to vector molecules, for introducing the composite DNA molecule into a host cell in which it can replicate, and for selecting the clone having the target gene from amongst the recipient host cells.

By "cDNA" is meant complementary or copy DNA produced from an RNA template by the action of RNA-dependent DNA polymerase (reverse transcriptase). Thus a "cDNA clone" means a duplex DNA sequence complementary to an RNA molecule of interest, carried in a cloning vector.

By "cDNA library" is meant a collection of recombinant DNA molecules containing cDNA inserts which together comprise a representation of the mRNA molecules present in an entire organism or tissue, depending on the source of the RNA templates. Such a cDNA library may be prepared by methods known to those of skill, and described, for example, in Maniatis et al., Molecular Cloning: A Laboratory Manual, supra. Generally, RNA is first isolated from the

cells of an organism from whose genome it is desired to clone a particular gene. Preferred for the purposes of the present invention are mammalian, and particularly human, cell lines.

Alternatively, RNA may be isolated from a tumor cell, derived from an animal tumor, and preferably from a human tumor. Thus, a library may be prepared from, for example, a human adrenal tumor, but any tumor may be used.

As used herein, the term "DNA polymorphism" refers to the condition in which two or more different nucleotide sequences can exist at a particular site in DNA.

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"Expression vector" includes vectors which are capable of expressing DNA sequences contained therein, i.e., the coding sequences are operably linked to other sequences capable of effecting their expression. It is implied, although not always explicitly stated, that these expression vectors must be replicable in the host organisms either as episomes or as an integral part of the chromosomal DNA. A useful, but not a necessary, element of an effective expression vector is a marker encoding sequence, which is a sequence encoding a protein which results in a phenotypic property (such as tetracycline resistance) of the cells containing the protein which permits those cells to be readily identified. In sum, "expression vector" is given a functional definition, and any DNA sequence which is capable of effecting expression of a specified contained DNA code is included in this term, as it is applied to the specified sequence. As at present, such vectors are frequently in the form of plasmids, thus "plasmid" and "expression vector" are often used interchangeably. However, the invention is intended to include such other forms of expression vectors which serve equivalent functions and which may, from time to time become known in the art.

By "functional derivative" is meant the "fragments", "variants", "analogs", or "chemical derivatives" of a molecule. A "fragment" of a molecule, such as any of the antigens of the present invention is meant to refer to any polypeptide subset of the molecule. A "variant" of such molecules is meant to refer to a naturally occurring molecule substantially similar to either the entire molecule, or a fragment thereof. An "analog" of a molecule is meant to refer to a non-natural molecule substantially similar to either the entire molecule or a fragment thereof.

The term "gene" means a polynucleotide sequence encoding a peptide.

By "homogeneous" is meant, when referring to a peptide or DNA sequence, that the primary molecular structure (i.e., the sequence of amino acids or nucleotides) of substantially all molecules present in the composition under consideration is identical.

"Isolated" refers to a protein of the present invention, or any gene encoding any such protein, which is essentially free of other proteins or genes, respectively, or of other contaminants with which it might normally be found in nature, and as such exists in a form not found in nature.

The term "label" refers to a molecular moiety capable of detection including, by way of example, without limitation, radioactive isotopes, enzymes, luminescent agents, and dyes.

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The term "probe" refers to a ligand of known qualities capable of selectively binding to a target antiligand. As applied to nucleic acids, the term "probe" refers to a strand of nucleic acid having a base sequence complementary to a target strand.

"Recombinant host cells" refers to cells which have been transformed with vectors constructed using recombinant DNA techniques. As defined herein, the antibody or modification thereof produced by a recombinant host cell is by virtue of this transformation, rather than in such lesser amounts, or more commonly, in such less than detectable amounts, as would be produced by the untransformed host.

By "substantially pure" is meant any protein of the present invention, or any gene encoding any such protein, which is essentially free of other proteins or genes, respectively, or of other contaminants with which it might normally be found in nature, and as such exists in a form not found in nature.

A molecule is said to be "substantially similar" to another molecule if the sequence of amino acids in both molecules is substantially the same, and if both molecules possess a similar biological activity. Thus, provided that two molecules possess a similar activity, they are considered variants as that term is used herein even if one of the molecules contains additional amino acid residues not found in the other, or if the sequence of amino acid residues is not identical. As used herein, a molecule is said to be a "chemical derivative" of another molecule when it contains additional chemical moieties not normally a part of the molecule. Such moieties may improve the molecule's solubility, absorption, biological half life, etc. The moieties may alternatively decrease the toxicity of the molecule, eliminate or attenuate any undesirable side effect of the molecule, etc. Moieties capable of mediating such effects are disclosed, for example, in Remington's Pharmaceutical Sciences, 16th ed., Mack Publishing Co., Easton, Penn. (1980).

- 10 -

By "vector" is meant a DNA molecule, derived from a plasmid or bacteriophage, into which fragments of DNA may be inserted or cloned. A vector will contain one or more unique restriction sites, and may be capable of autonomous replication in a defined host or vehicle organism such that the cloned sequence is reproducible.

5 Compounds of the Invention

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The invention includes the cDNA of SEQ ID NO: 1, SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6, as well as sequences which include the sequence of SEQ ID NO: 1, SEQ ID NO:2, SEQ ID NO:4, or SEQ ID NO:6, and derivatives of these sequences. The invention also includes vectors, liposomes and other carrier vehicles which encompass these sequence or derivatives of them. The invention further includes proteins transcribed from SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:4, or SEQ ID NO:6, including but not limited to SEQ ID NO:3, SEQ ID NO:5 or SEQ ID NO:7, and their derivatives and variants.

One embodiment of the invention includes soluble variants of a KIM protein that is usually synthesized as a membrane associated protein, and which is upregulated after injury. Soluble variants lack at least a portion of the transmembrane or intra-membrane section of a native KIM protein. In some examples, the soluble variant lacks the entire transmembrane or intra-membrane section of a native KIM protein. Soluble variants include fusion proteins which encompass derivatives of KIM proteins that lack at least a portion of the transmembrane or intra-membrane section of a native KIM protein. All types of KIM fusion proteins are included, particularly those which incorporate his-tag, Ig-tag, and myc-tag forms of the molecule. These KIM fusions may have characteristics which are therapeutically advantageous, such as the increased half-life conferred by the Ig-tag. Also included are fusion proteins which incorporate portions of selected domains of the KIM protein.

Variants can differ from naturally occurring KIM protein in amino acid sequence or in ways that do not involve sequence, or both. Variants in amino acid sequence are produced when one or more amino acids in naturally occurring KIM protein is substituted with a different natural amino acid, an amino acid derivative or non-native amino acid. Particularly preferred variants include naturally occurring KIM protein, or biologically active fragments of naturally occurring KIM protein, whose sequences differ from the wild type sequence by one or more conservative amino acid substitutions, which typically have minimal influence on the secondary structure and

- 11 -

hydrophobic nature of the protein or peptide. Variants may also have sequences which differ by one or more non-conservative amino acid substitutions, deletions or insertions which do not abolish the KIM protein biological activity. Conservative substitutions typically include the substitution of one amino acid for another with similar characteristics such as substitutions within the following groups: valine, glycine; glycine, alanine; valine, isoleucine; aspartic acid, glutamic acid; asparagine, glutamine; serine, threonine; lysine, arginine; and phenylalanine, tyrosine. The non-polar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan and methionine. The polar neutral amino acids include glycine, serine, threonine, cysteine, tyrosine, asparagine and glutamine. The positively charged (basic) amino acids include arginine, lysine and histidine. The negatively charged (acidic) amino acids include aspartic acid and glutamic acid.

Other conservative substitutions can be taken from the table below, and yet others are described by Dayhoff in the Atlas of Protein Sequence and Structure (1988).

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- 12 -

TABLE 1: CONSERVATIVE AMINO ACID REPLACEMENTS

For Amino	Acid	Code	Replace with any of				
Alanine		A	D-Ala, Gly,beta-Ala, L- Cys,D-Cys				
Arginine		R	D-Arg, Lys,homo-Arg, D-homo-Arg, Met,D-Met, Ile, D-Ile, Orn, D-Orn				
5 Asparagine		N	D-Asn,Asp,D-Asp,Glu,D-Glu, Gln,D-Gln				
Aspartic Ac	id	D	D-Asp,D-Asn,Asn, Glu,D-Glu, Gln, D-Gln				
Cysteine		С	D-Cys, S-Me-Cys,Met,D- Met,Thr, D-Thr				
Glutamine		Q	D-Gln,Asn, D-Asn,Glu,D- Glu,Asp, D-Asp				
Glutamic A	cid	Е	D-Glu,D-Asp,Asp, Asn, D-Asn, Gln, D-Gln				
) Glycine		G	Ala, D-Ala, Pro, D-Pro, Beta- Ala, Acp				
Isoleucine		I	D-Ile, Val, D-Val, Leu, D- Leu, Met, D-Met				
Leucine		L,	D-Leu, Val, D-Val, Met, D-Met				
Lysine		K	D-Lys,Arg, D-Arg, homo-Arg, D-homo-Arg, Met, D_Met, Ile, D-Ile, Orn, D-Orn				
Methionine	:	М	D-Met, S-Me-Cys, Ile, D-Ile, Leu, D-Leu, Val, D-Val, Norieu				
5 Phenylalan	ine	F	D-Phe,Tyr, D-Thr,L- Dopa,His,D-His, Trp, D-Trp, Trans 3,4 or 5-phenylproline, cis 3,4 or 5 phenylproline				

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- 13 -

Proline	P	D-Pro, L-I-thioazolidine-4- carboxylic acid, D- or L-1- oxazolidine-4-carboxylic acid
Serine	S	D-Ser, Thr, D-Thr, allo-Thr, Met, D-Met, Met(O), D-Met(O), Val, D-Val
Threonine	Т	D-Thr, Ser, D-Ser, allo-Thr, Met, D-Met, Met)O, D- Met(O), Val, D-Val
Tyrosine	Y	D-Tyr,Phe, D-Phe, L-Dopa, His,D-His
Valine	V	D-Val, Leu, D-Leu, Ile, D-Ile, Met, D-Met

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- 14 -

Other variants within the invention are those with modifications which increase peptide stability. Such variants may contain, for example, one or more non-peptide bonds (which replace the peptide bonds) in the peptide sequence. Also included are: variants that include residues other than naturally occurring L-amino acids, such as D-amino acids or non-naturally occurring or synthetic amino acids such as beta or gamma amino acids and cyclic variants. Incorporation of D- instead of L-amino acids into the polypeptide may increase its resistance to proteases. See, e.g., U.S. Patent 5,219,990.

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Generally, substitutions that may be expected to induce changes in the functional properties of KIM polypeptides are those in which: (I) a hydrophilic residue, e.g., serine or threonine, is substituted by a hydrophobic residue, e.g., leucine, isoleucine, phenylalanine, or alanine; (ii) a cysteine residue is substituted for (or by) any other residue; (iii) a residue having an electropositive side chain, e.g., lysine, arginine or histidine, is substituted for (or by) a residue having an electronegative charge, e.g., glutamic acid or aspartic acid; or (iv) a residue having a bulky side chain, e.g., phenylalanine, is substituted for (or by) one not having such a side chain, e.g., glycine.

The peptides of this invention may also be modified by various changes such as insertions, deletions and substitutions, either conservative or nonconservative where such changes might provide for certain advantages in their use. Splice variants are specifically included in the invention.

In other embodiments, variants with amino acid substitutions which are less conservative may also result in desired derivatives, e.g., by causing changes in charge, conformation and other biological properties. Such substitutions would include for example, substitution of hydrophilic residue for a hydrophobic residue, substitution of a cysteine or proline for another residue, substitution of a residue having a small side chain for a residue having a bulky side chain or substitution of a residue having a net positive charge for a residue having a net negative charge. When the result of a given substitution cannot be predicted with certainty, the derivatives may be readily assayed according to the methods disclosed herein to determine the presence or absence of the desired characteristics.

Variants within the scope of the invention include proteins and peptides with amino acid sequences having at least eighty percent homology with a KIM protein. More preferably the sequence homology is at least ninety percent, or at least ninety-five percent. For the purposes

- 15 -

of determining homology the length of comparison sequences will generally be at least 8 amino acid residues, usually at least 20 amino acid residues. Variants of the compounds of the invention also includes any protein which 1) has an amino acid sequence which is at least forty percent homologous to a KIM protein of the invention, and also which 2) after being placed in an optimal alignment with the KIM sequence (as depicted in Figure 5 for human and for rat KIM-1) has at least 80% of its cysteine residues aligned with cysteines in the KIM protein of the invention.

Just as it is possible to replace substituents of the scaffold, it is also possible to substitute functional groups which are bound to the scaffold with groups characterized by similar features. These substitutions will initially be conservative, i.e., the replacement group will have approximately the same size, shape, hydrophobicity and charge as the original group. Non-sequence modifications may include, for example, *in vivo* or *in vitro* chemical derivatization of portions of naturally occurring KIM protein, as well as changes in acetylation, methylation, phosphorylation, carboxylation or glycosylation.

Also included within the invention are agents which specifically bind to the protein, or a fragment of the protein (SEQ ID NO:3, 5 or 7). These agents include ligands and antibodies (including monoclonal, single chain, double chain, Fab fragments, and others, whether native, human, humanized, primatized, or chimeric). Additional descriptions of these categories of agents are in PCT application 95/16709, the specification of which is herein incorporated by reference.

Experimental Procedures

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1. Generation of RNA from ischemic and normal rat adult kidneys

Ischemic injured rat kidneys are generated as described by Witzgall et al. (J. Clin Invest. 93: 2175-2188, 1994). Briefly, the renal artery and vein from one kidney of an adult Sprague-Dawley rat are clamped for 40 minutes and then reperfused. Injured kidneys are harvested from the rats at 24 hours and at 48 hours after reperfusion. Kidneys from sham-operated, normal adult Sprague-Dawley rats are also harvested.

Total RNA is prepared from the organs based on the protocol by Glisin et al.

(Biochemistry 13: 2633, 1974). Briefly, the harvested organs are placed immediately into GNC

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buffer (4M guanidine thiocyanate, 0.5% SDS, 25mM sodium citrate, 0.1% Sigma anti foam) and disrupted on ice with a polytron. Cell debris is removed with a low speed spin in a clinical centrifuge and the supernatant fluid is placed on a 5.7 M CsCl, 25mM sodium acetate, 1mM EDTA cushion. RNA is pelleted through the cushion in a SW40Ti rotor at 22K for 15hrs. RNA is resuspended in sterile DEPC- treated water, precipitated twice with 1/10 volume 3M sodium acetate and 2.5 volumes of EtOH. Poly A+ RNA is isolated using an mRNA purification kit (Pharmacia, catalog No.27-9258-02).

2. Representational Difference Analysis (RDA) method to isolate 1-7. 3-2 and 4-7 RDA fragments

Double stranded cDNA is synthesized from sham-operated and from 48hr post-ischemic kidney poly A+ RNA using Gibco BRL "Superscript Choice™ System cDNA Synthesis Kit", catalog No. 18090. First strand is synthesized by priming with oligo dT and using Superscript II™ reverse transcriptase. Second strand is generated using E. coli DNA polymerase I and RNase H followed by T4 DNA polymerase using BRL recommended conditions.

RDA analysis is performed essentially as described by Hubank and Schatz (Nucleic Acid Research 22: 5640-48, 1994). Briefly, 48 hr post-ischemic kidney cDNA is digested with the restriction enzyme *Dpn* II, and ligated to R-Bgl-12/24 oligonucleotides (see reference for exact sequence). PCR amplification (performed with Perkin-Elmer Taq polymerase and their corresponding PCR buffer) of the linker ligated cDNA is used to generate the initial representation. This PCR product is designated "tester amplicon." The same procedure is used to generate "driver amplicon" from sham-operated rat kidney cDNA.

Hybridization of tester and driver amplicons followed by selective amplification are performed three times to generate Differential Product One (DP1), Two (DP2) and Three (DP3). Generation of the DP1 product is performed as described by Hubank and Schatz (Nucleic Acid Research 22: 5640-48, 1994). The DP2 and DP3 products are also generated as described by Hubank and Schatz (id.), except that the driver:tester ratios are changed to 5,333:1 for DP2 and to 40,000:1 or 4,000:1 for DP3.

Three RDA products are cloned from DP3 into the cloning vector pUC 18: RDA product 1-7 (252bp) when the DP3 was generated using a ratio of 40,000:1, and product RDA 3-2 (445bp) and 4-7 (483bp) when the DP3 was generated using a ratio of 4,000:1. The DNA

- 17 -

fragments are subcloned using the Pharmacia Sureclone™ kit (catalog No. 27-9300-01) to repair the ends of the PCR fragments with Klenow enzyme and to facilitate blunt end ligation of the fragments into the pUC18 vector.

3. Northern Analysis

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Poly A+ RNA (2.5μg) from rat normal adult kidney (sham operated), from 48hr post-ischemic injured adult kidney, and from day 18 embryonic kidney is electrophoresed and Northern blotted (Cate, Cell 45:685, 1986) to a GeneScreenTM membrane (Dupont).

Hybridization in PSB buffer (50mM Tris 7.5, 1M NaCl, 0.1% Na pyrophosphate, 0.2% PVP, 0.2% Ficoll, 0.2% BSA, 1% SDS), containing 10% dextran sulphate and 100μg/ml tRNA, is performed at 65C using three different probes: 1-7 RDA product, 3-2 RDA product and 4-7 RDA product. All are radiolabeled using Pharmacia's "Ready to GoTM" random priming labeling kit (catalog No.27-9251-01). RDA products 1-7, 3-2 and 4-7 hybridize to mRNAs present in all three samples, but most intensely to mRNAs in the 48hr post-ischemic kidney RNA samples.

A Northern blot analysis of adult rat tissues indicates that the 1-7 gene is expressed at very low levels in normal adult kidney, testis, spleen and lung. The 3-2 gene is expressed in liver, kidney, spleen, and brain. The 4-7 gene is expressed in spleen, kidney, lung, testis, heart, brain, liver, and skeletal muscle. The presence of different sized mRNAs in some tissues in the 1-7 and 3-2 blot indicates that the primary transcription product of the 1-7 gene and of the 3-2 gene may undergo alternate splicing and/or polyadenylation.

20 4. Isolation of 3-2 and 4-7 cDNA clones

A cDNA library is generated from 4 μg of polyA+ RNA from 48hr post-ischemic injured kidney using reagents from BRL Superscript ChoiceTM System for cDNA synthesis, and StratageneTM Lambda ZapII cloning kit (catalog No. 236201), according to protocols recommended by the manufacturers.

25 10³ clones are screened with the 3-2 RDA product as a probe (random primed labeled as described above). Eight positive clones are selected and four are randomly chosen for secondary analysis to obtain pure phage plaques. After tertiary screening, four pure phage clones are isolated. Cloned inserts from the phage are isolated by in vivo excision procedure according to Stratagene™ Lambda Zap II kit. The largest insert, of approximately 2.6 kb (referred to as

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cDNA clone 3-2), is subjected to DNA sequencing. The sequence of the insert (SEQ ID NO:1) is shown in Figure 1. cDNA clone 3-2 (*E. coli* K-12, SOLR/p3-2#5-1) has been deposited as ATCC No. 98061. The sequence of cDNA clone 3-2 is identical to that of clone 1-7 cDNA (SEQ ID NO: 2), except that nucleotides 136-605 of SEQ ID NO:1 represent an insertion. Thus, SEQ ID NO:2 represents a splice variant form of SEQ ID NO: 1. The clone for 1-7 (*E. coli* K-12, SOLR/p1-7#3-1) has been deposited as ATCC No. 98060.

10⁵ clones are screened with the 1-7 RDA product as a probe (random primed radiolabeled as described above). Eight positive clones are selected and four are randomly chosen for secondary analysis to obtain pure phage plaques. After tertiary screening, four pure phage clones are isolated. Cloned inserts from the phage are isolated by *in vivo* excision procedure according to Stratagene™ Lambda Zap II kit. The largest insert of approximately 2.0 kb (referred to as cDNA clone 1-7) is subjected to DNA sequencing; the sequence of the insert (SEO ID NO: 2) is shown in Figure 2.

10⁵ clones are screened with the 4-7 RDA product as a probe (random primed labeled as described above and hybridized in PSB at 65C). Eight positive clones are selected and four are randomly chosen for secondary analysis to obtain pure phage plaques. After secondary screening, two pure phage clones are isolated. Cloned inserts from the phage are isolated by *in vivo* excision procedure according to Stratagene™ Lambda Zap II kit. The largest insert, approximately 2.4 kb (referred to as cDNA clone 4-7), is subjected to DNA sequencing. The sequence of the insert, SEQ ID NO: 4, is shown in Figure 3. The cDNA clone 4-7 (*E. coli* K-12, SOLR/p4-7#1-1) has been deposited as ATCC No. 98062..

5. Characterization of the 1-7. 3-2 and 4-7 cDNA clones

A.) DNA and Protein Sequences:

The sequence of 3-2 cDNA (Figure 1; SEQ ID NO:1) contains an open reading frame of 307 amino acids (Figure 1; SEQ ID NO:3). A signal sequence of 21 amino acids is inferred from Von Heijne analysis (Von Heijne et al., Nucl. Acid Res. 14:14683 (1986)), and a transmembrane region spanning approximately aa 235-257 indicates that the 3-2 product is a cell surface protein.

The sequence of 1-7 cDNA (Figure 2; SEQ ID NO:2) contains an open reading frame of 307 amino acids, which is identical to the open reading frame contained in the 3-2 cDNA (SEQ ID NO: 3). The sequence of 4-7 cDNA (Figure 3; SEQ ID NO:4) contains an open reading

frame of 572 amino acids (SEQ ID NO:5). A transmembrane region is located at approximately amino acids 501-521.

B.) In situ analysis of 1-7, 3-2 and 4-7 mRNAs in contralateral and in post-ischemic adult rat kidneys:

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In situ hybridization is carried out according to the method described by Finch et al., Dev. Dynamics 203: 223-240, 1995. Briefly, both ischemic and contralateral kidneys are perfusion fixed with 4% paraformaldehyde in PBS. Kidneys are further fixed overnight at 4C and processed. Paraffin sections are deparaffinized and rehydrated, fixed with 4% paraformaldehyde in PBS, digested with proteinase K, refixed, then acetylated with acetic anhydride in triethanolamine buffer. Sections are then dehydrated and hybridized with ³²P-labeled riboprobes at 55°C overnight, with 33P-labeled riboprobes generated from 3-2 RDA or 1-7 RDA products subcloned into BamH1 site of pGEM-11Z. After hybridization, sections were washed under high stringency conditions (2 X SSC, 50 % formamide at 65°C). Sections are finally dehydrated, emulsion (NBT-2) coated for autoradiography, and exposed for at least a week. Silver grains are developed and sections are counterstained with toluidine blue and microphotographed.

Analysis of 1-7 and 3-2 mRNA expression by *in situ* hybridization indicates that these genes are greatly upregulated in damaged kidney cells compared to their expression in normal kidney sections. The expression seen is in regenerative cells of the cortex and outer medulla, most of which appear to be proximal tubule cells.

Analysis of the 4-7 in situ RNA expression pattern also reveals abundant expression of this gene in the injured ischemic kidney compared to the normal adult kidney. The site of expression appears to be infiltrating cells.

6.) Isolation of a human cDNA clone which cross hybridizes to the rat 3-2 cDNA

A ³²P-labeled DNA probe comprising nucleotides 546-969 of the insert of clone 3-2 shown in Figure 1 is generated and used to screen a human embryonic liver lambda gt10 cDNA library (Clontech Catalog #HL5003a). 1 X10⁶ plaques are screened in duplicate using standard conditions as described above but temperature for screening was 55C. For the high stringency wash, the filters are washed in 2X SSC at 55C. Fifty positive phage are identified and plaque

- 20 -

purified, and DNA is prepared. The phage DNAs are subjected to Southern analysis using the same probe as above. The Southern blot filter is subjected to a final wash with 0.5X SSC at 55C. Two clones are identified as positive. The insert of clone H13-10-85 is sequenced and a region is found that encodes a protein with a high level of identity to the 3-2 protein shown in Figure 3.

The nucleotide sequence (SEQ ID NO:6) and predicted amino acid sequence (SEQ ID NO:7) of the human 3-2 related protein are shown in Figure 4. As shown by the bestfit analysis depicted in Figure 5, the human 3-2 related protein is 43.8% identical and 59.1% similar to the rat 3-2 protein. Both contain IgG, mucin, transmembrane, and cytoplasmic domains. The six cysteines within the IgG domains of both proteins are conserved.

7) Production of KIM-1 Ig fusion protein

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A fusion protein of the extracellular domain of KIM and the Fc region of immunoglobulin (Ig) is a useful tool for the study of the molecular and cellular biology of the injured/regenerating kidney and as a therapeutic molecule. To produce Kim Ig fusion protein with the extracellular domain of human and rat KIM-1 protein, a fragment of the extracellular domain of KIM-1 cDNA was amplified by PCR and cloned in the Biogen expression vector, pCA125, for transient expression in COS cells. The expression vector pCA125 produces a fusion protein which has a structure from gene cloned at N-terminus and a human Ig Fc region at the C-terminus. COS cells were transfected with the plasmids SJR 103 or 104; these plasmids express a fusion protein which contains the human KIM sequences 263-1147 (SEQ ID NO:6; SJR 103) or rat KIM sequences 599-1319 (SEQ ID NO:1; SJR 104) of the extracellular domain fused to human Ig Fc region. The cells were grown in 10% FBS in DMEM in the cell factory (Nunc, Naperville, II).Two to three days post-transfection, medium was harvested, concentrated using Amicon concentrator, and fusion protein was purified using Protein-A Sepharose column. After purification, purity of fusion protein was evaluated by SDS-PAGE.

25 Diagnostic Uses of the Compounds of the Invention

Anti-KIM antibodies of the invention, which specifically bind to the protein of SEQ ID NO:3, SEQ ID NO:5 or SEQ ID NO:7 or a fragment thereof, are useful in several diagnostic methods. These agents may be labeled with detectable markers, such as fluoroscopically or radiographically opaque substances, and administered to a subject to allow imaging of tissues

-21 -

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which express KIM protein. The agents may also be bound to substances, such as horseradish peroxidase, which can be used as immunocytochemical stains to allow visualization of areas of KIM protein-positive cells on histological sections. A specific antibody could be used alone in this manner, and sites where it is bound can be visualized in a sandwich assay using an anti-immunoglobulin antibody which is itself bound to a detectable marker.

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Specific antibodies to KIM protein are also useful in immunoassays to measure KIM presence or concentration in samples of body tissues and fluids. Such concentrations may be correlated with different disease states. As an embodiment of particular interest, the invention includes a method of diagnosing renal injury, or of monitoring a process of renal repair, by measuring the concentration of KIM or of KIM fragments in the urine, plasma or serum of a patient. Similarly, KIM can be measured in urine sediment, in particular in cellular debris in the urine sediment. Casts of renal tubule cells, which may be present in urine sediment from patients with ongoing renal disease, may contain elevated levels of KIM protein and mRNA.

Specific antibodies to KIM protein may also be bound to solid supports, such as beads or dishes, and used to remove the ligand from a solution, either for measurement, or for purification and characterization of the protein or its attributes (such as posttranslational modifications). Such characterization of a patient's KIM protein might be useful in identifying deleterious mutants or processing defects which interfere with KIM function and are associated with abnormal patient phenotypes. Each of these techniques is routine to those of skill in the immunological arts.

Additional imaging methods utilize KIM or KIM fragments, fused to imageable moieties, for diagnostic imaging of tissues that express KIM ligands, particularly tumors.

Further diagnostic techniques are based on demonstration of upregulated KIM mRNA in tissues, as an indication of injury-related processes. This technique has been tested and found workable in a model of ischemic injury in rats, as follows.

To determine if the amount of KIM-1 protein is increased after injury, we examined kidney homogenates of contralateral and postischemic kidneys 24 and 48 hours following a 40 minute clamping of the renal artery and vein of a single kidney for each rat. The kidney homogenate was assessed for the presence of KIM-1 protein. Western blot analysis identifies three proteins detected by two different antibodies after ischemic injury, which are not detectable in homogenates from contralateral kidneys which were not exposed to ischemic injury. The

apparent molecular weights of the bands are approximately 40kDa, 50kDa and 70-80kDa. The three protein species detected by western blotting could represent glycosylated forms of the same protein given the presence of potential N and O linked glycosylation sites. The fact that each of these proteins react with two different sets of polyclonal antibodies supports the idea that they are related to KIM-1 and are not cross-reacting bands. Confirmation of this prediction came from the results of partial CNBr cleavage of the three proteins which revealed they shared common CNBr cleavage fragments. Since the cytoplasmic domain of the KIM-1 protein is not predicted to contain any major post-translational modifications, the two smallest products of the digest (4.7kDa and 7.4kDa) detected with antibodies directed against the cytoplasmic domain of KIM-1 should be the same size for the three different KIM-1 protein bands if they originate from the same protein. We observed that the KIM1 40kDa and 70-80kDa proteins yield fragments migrating at the predicted size. Digest of the 50kDa protein band gave also the same C-terminal signature band peptide.

The KIM-1 sequence presents two putative sites for N-glycosylation and a mucin domain where O-glycosylation could cover the polypeptide chain. The three KIM-1 bands detected in postischemic kidney could correspond to glycosylation variants of the same core protein. De-N-glycosylation with PNGase F resulted in a shift of all three bands to a lower molecular weight, corresponding to a loss of about 3kDa, indicating that all three proteins are N-glycosylated. Differences in O-glycosylation might explain the differences in sizes of these three bands.

20 Therapeutic Uses of the Compounds of the Invention

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The therapeutic methods of the invention involve selectively promoting or inhibiting cellular responses that are dependent on KIM ligation. Where the KIM and the KIM ligand are both membrane bound, and expressed by different cells, the signal transduction may occur in the KIM-expressing cell, in the KIM ligand-expressing cell, or in both.

KIM ligation-triggered response in a KIM ligand-expressing cell may be generated by contacting the cell with exogenous KIM, KIM fusion proteins or activating antibodies against KIM ligand, either in vitro or in vivo. Further, responses of the KIM ligand-expressing cell that would otherwise be triggered by endogenous KIM could be blocked by contacting the KIM ligand-expressing cell with a KIM ligand antagonist (e.g., an antagonist antibody that binds to

- 23 -

KIM ligand), or by contacting the endogenous KIM with an anti-KIM antibody or other KIM-binding molecule which prevents the effective ligation of KIM with a KIM ligand.

Similarly, the responses triggered by KIM ligation in the KIM-expressing cell may be promoted or inhibited with exogenous compounds. For example, KIM ligation-triggered response in a KIM-expressing cell may be generated by contacting the cell with a soluble KIM ligand, or certain anti-KIM activating antibodies. Further, responses of the KIM-expressing cell that would otherwise be triggered by interaction with endogenous KIM ligand could be blocked by contacting the KIM-expressing cell with an antagonist to KIM (e.g.., a blocking antibody that binds to KIM in a manner that prevents effective, signal-generating KIM ligation), or by contacting the endogenous KIM ligand with an anti-KIM ligand antibody or other KIM ligand-binding molecule which prevents the effective ligation of KIM with the KIM ligand.

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Which of the interventions described above are useful for particular therapeutic uses depend on the relevant etiologic mechanism of either the pathologic process to be inhibited, or of the medically desirable process to be promoted, as is apparent to those of skill in the medical arts. For example, where KIM ligation results in desirable cellular growth, maintenance of differentiated phenotype, resistance to apoptosis induced by various insults, or other medically advantageous responses, one of the above-described interventions that promote ligation-triggered response may be employed. In the alternative, one of the inhibitory interventions may be useful where KIM ligation invokes undesirable consequences, such as neoplastic growth, deleterious loss of cellular function, susceptibility to apoptosis, or promotion of inflammation events.

Following are examples of the previously described therapeutic methods of the invention. One therapeutic use of the KIM-related compounds of the invention is for treating a subject with renal disease, promoting growth of new tissue in a subject, or promoting survival of damaged tissue in a subject, and includes the step of administering to the subject a therapeutically effective amount of a KIM protein of the invention, or of a pharmaceutical composition which includes a protein of the invention. The protein used in these methods may be a fragment of a full-length KIM protein, a soluble KIM ligand protein or fusion fragment, or a KIM agonist. These methods may also be practiced by administering to the subject a therapeutically effective amount of an agonist antibody of the invention, or a pharmaceutical composition which includes an agonist antibody of the invention. A KIM protein may be administered concurrently with a therapeutically effective amount of a second compound which exerts a medically desirable

- 24 -

adjunct effect. While tissues of interest for these methods may include any tissue, preferred tissues include renal tissue, liver, neural tissue, heart, stomach, small intestine, spinal cord, or lung. Particular renal conditions which may be beneficially treated with the compounds of the invention include acute renal failure, acute nephritis, chronic renal failure, nephrotic syndrome, renal tubule defects, kidney transplants, toxic injury, hypoxic injury, and trauma. Renal tubule defects include those of either hereditary or acquired nature, such as polycystic renal disease, medullary cystic disease, and medullary sponge kidney. This list is not limited, and may include many other renal disorders (see, e.g., Harrison's Principles of Internal Medicine, 13th ed., 1994, which is herein incorporated by reference.) The subject of the methods may be human.

A therapeutic intervention for inhibiting growth of undesirable, KIM ligand-expressing tissue in a subject includes the step of administering to the subject a therapeutically effective amount of a KIM antagonist (e.g.., an antagonist antibody that binds to KIM ligand), or by administering a therapeutically effective amount of an anti-KIM antibody or other KIM-binding molecule which blocks KIM binding to the KIM ligand-expressing tissue. In an embodiment of interest, the KIM antagonist or anti-KIM antibody may be used therapeutically to inhibit or block growth of tumors which depend on KIM protein for growth.

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Other methods of the invention include killing KIM ligand-expressing tumor cells, or inhibiting their growth, by contacting the cells with a fusion protein of a KIM and a toxin or radionuclide, or an anti-KIM ligand antibody conjugated to a toxin or radionuclide. The cell may be within a subject, and the protein or the conjugated antibody is administered to the subject.

Also encompassed within the invention is a method for targeting a toxin or radionuclide to a cell expressing a KIM, comprising contacting the cell with a fusion protein comprising a KIM ligand and a toxin or radionuclide, or an anti-KIM antibody conjugated to a toxin or radionuclide. Another embodiment includes the method of suppressing growth of a tumor cell which expresses KIM, comprising contacting the cell with a fusion protein of KIM ligand and a toxin or radionuclide or with an anti-KIM antibody conjugated to a toxin or radionuclide; the cell may be within a subject, and the protein administered to the subject.

The term "subject" used herein is taken to mean any mammal to which KIM may be administered. Subjects specifically intended for treatment with the method of the invention include humans, as well as nonhuman primates, sheep, horses, cattle, goats, pigs, dogs, cats,

rabbits, guinea pigs, hamsters, gerbils, rats and mice, as well as the organs, tumors, and cells derived or originating from these hosts.

Use of Compounds of the Invention in Gene Therapy

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The KIM genes of the invention are introduced into damaged tissue, or into tissue where stimulated growth is desirable. Such gene therapy stimulates production of KIM protein by the transfected cells, promoting cell growth and/or survival of cells that express the KIM protein.

In a specific embodiment of a gene therapy method, a gene coding for a KIM protein may be introduced into a renal target tissue. The KIM protein would be stably expressed and stimulate tissue growth, division, or differentiation, or could potentiate cell survival.

Furthermore, a KIM gene may be introduced into a target cell using a variety of well-known methods that use either viral or non-viral based strategies.

Non-viral methods include electroporation, membrane fusion with liposomes, high velocity bombardment with DNA-coated microprojectiles, incubation with calcium-phosphate-DNA precipitate, DEAE-dextran mediated transfection, and direct micro-injection into single cells. For instance, a KIM gene may be introduced into a cell by calcium phosphate coprecipitation (Pillicer et al., Science, 209: 1414-1422 (1980); mechanical microinjection and/or particle acceleration (Anderson et al., Proc. Nat. Acad. Sci. USA, 77: 5399-5403 (1980); liposome based DNA transfer (e.g., LIPOFECTIN-mediated transfection- Fefgner et al., Proc. Nat. Acad. Sci., USA, 84: 471-477, 1987; Gao and Huang, Biochim. Biophys. Res. Comm., 179: 280-285, 1991; DEAE Dextran-mediated transfection; electroporation (U.S. Patent 4,956,288); or polylysine-based methods in which DNA is conjugated to deliver DNA preferentially to liver hepatocytes (Wolff et al., Science, 247: 465-468, 1990; Curiel et al., Human Gene Therapy 3: 147-154, 1992).

Target cells may be transfected with the genes of the invention by direct gene transfer.

See, e.g., Wolff et al., "Direct Gene Transfer Into Moose Muscle In Vivo", Science 247:1465-68, 1990. In many cases, vector-mediated transfection will be desirable. Any of the methods known in the art for the insertion of polynucleotide sequences into a vector may be used. (See, for example, Sambrook et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY, 1989; and Ausubel et al., Current Protocols in Molecular Biology, J. Wiley & Sons, NY, 1992, both of which are incorporated herein by reference.)

Promoter activation may be tissue specific or inducible by a metabolic product or administered substance. Such promoters/enhancers include, but are not limited to, the native c-ret ligand protein promoter, the cytomegalovirus immediate-early promoter/enhancer (Karasuyama et al., J. Exp. Med., 169: 13, 1989); the human beta-actin promoter (Gunning et al., Proc. Nat. Acad. Sci. USA, 84: 4831, 1987; the glucocorticoid-inducible promoter present in the mouse mammary tumor virus long terminal repeat (MMTV LTR) (Klessig et al., Mol. Cell. Biol., 4: 1354, 1984); the long terminal repeat sequences of Moloney murine leukemia virus (MuLV LTR) (Weiss et al., RNA Tumor Viruses, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY, 1985); the SV40 early region promoter (Bernoist and Chambon, Nature, 290:304, 1981); the promoter of the Rous sarcoma virus (RSV) (Yamamoto et al., Cell, 22:787, 1980); the herpes simplex virus (HSV) thymidine kinase promoter (Wagner et al., Proc. Nat. Acad. Sci. USA, 78: 1441, 1981); the adenovirus promoter (Yamada et al., Proc. Nat. Acad. Sci. USA, 82: 3567, 1985).

The KIM genes may also be introduced by specific viral vectors for use in gene transfer systems which are now well established. See for example: Madzak et al., J. Gen. Virol., 73: 1533-36, 1992 (papovavirus SV40); Berkner et al., Curr. Top. Microbiol. Immunol., 158: 39-61, 1992 (adenovirus); Hofmann et al., Proc. Natl. Acad. Sci. 92: 10099-10103, 1995 (baculovirus); Moss et al., Curr. Top. Microbiol. Immunol., 158: 25-38, 1992 (vaccinia virus); Muzyczka, Curr. Top. Microbiol. Immunol., 158: 97-123, 1992 (adeno-associated virus); Margulskee, Curr. Top. Microbiol. Immunol., 158: 67-93, 1992 (herpes simplex virus (HSV) and Epstein-Barr virus (HBV)); Miller, Curr. Top. Microbiol. Immunol., 158: 1-24, 1992 (retrovirus); Brandyopadhyay et al., Mol. Cell. Biol., 4: 749-754, 1984 (retrovirus); Miller et al., Nature, 357: 455-450, 1992 (retrovirus); Anderson, Science, 256: 808-813, 1992 (retrovirus), Current Protocols in Molecular Biology: Sections 9.10-9.14 (Ausubel et al., Eds.), Greene Publishing Associcates, 1989, all of which are incorporated herein by reference.

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Preferred vectors are DNA viruses that include adenoviruses (preferably Ad-2 or Ad-5 based vectors), baculovirus, herpes viruses (preferably herpes simplex virus based vectors), and parvoviruses (preferably "defective" or non-autonomous parvovirus based vectors, more preferably adeno-associated virus based vectors, most preferably AAV-2 based vectors). See, e.g., Ali et al., Gene Therapy 1: 367-384, 1994; U.S. Patent 4,797,368 and 5,399,346 and discussion below.

- 27 -

The choice of a particular vector system for transferring, for instance, a KIM sequence will depend on a variety of factors. One important factor is the nature of the target cell population. Although retroviral vectors have been extensively studied and used in a number of gene therapy applications, they are generally unsuited for infecting cells that are not dividing but may be useful in cancer therapy since they only integrate and express their genes in replicating cells. They are useful for ex vivo approaches and are attractive in this regard due to their stable integration into the target cell genome.

Adenoviruses are eukaryotic DNA viruses that can be modified to efficiently deliver a

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therapeutic or reporter transgene to a variety of cell types. The general adenoviruses types 2 and 5 (Ad2 and Ad5, respectively), which cause respiratory disease in humans, are currently being developed for gene therapy of Duchenne Muscular Dystrophy (DMD)and Cystic Fibrosis (CF). Both Ad2 and Ad5 belong to a subclass of adenovirus that are not associated with human malignancies. Adenovirus vectors are capable of providing extremely high levels of transgene delivery to virtually all cell types, regardless of the mitotic state. High titers (1013 plaque forming units/ml) of recombinant virus can be easily generated in 293 cells (an adenovirustransformed, complementation human embryonic kidney cell line: ATCC CRL1573) and cryostored for extended periods without appreciable losses. The efficacy of this system in delivering a therapeutic transgene in vivo that complements a genetic imbalance has been demonstrated in animal models of various disorders. See Watanabe, Atherosclerosis, 36: 261-268, 1986; Tanzawa et al., FEBS Letters 118(1):81-84, 1980; Golasten et al., New Engl.J. Med. 309:288-296, 1983; Ishibashi et al., J. Clin. Invest. 92: 883-893, 1993; and Ishibashi et al., J. Clin. Invest. 93: 1889-1893, 1994, all of which are incorporated herein by reference. Indeed, recombinant replication defective adenovirus encoding a cDNA for the cystic fibrosis transmembrane regulator (CFTR) has been approved for use in at least two human CF clinical trials. See, e.g., Wilson, Nature 365:691-692, 1993. Further support of the safety of recombinant adenoviruses for gene therapy is the extensive experience of live adenovirus vaccines in human populations.

The first-generation recombinant, replication-deficient adenoviruses which have been developed for gene therapy of DMD and other inherited disorders contain deletions of the entire E1a and part of the E1b regions. This replication-defective virus is grown in 293 cells containing a functional adenovirus E1a gene which provides a transacting E1a protein. E1-deleted viruses are capable of replicating and producing infectious virus in the 293 cells, which provide E1a and

- 28 -

Elb region gene products in *trans*. The resulting virus is capable of infecting many cell types and can express the introduced gene (providing it carries its own promoter), but cannot replicate in a cell that does not carry the El region DNA unless the cell is infected at a very high multiplicity of infection. Adenoviruses have the advantage that they have a broad host range, can infect quiescent or terminally differentiated cells such as neurons, and appear essentially non-oncogenic. Adenoviruses do not appear to integrate into the host genome. Because they exist extrachromasomally, the risk of insertional mutagenesis is greatly reduced. Ali et al., supra, at 373. Recombinant adenoviruses (rAdV) produce very high titers, the viral particles are moderately stable, expression levels are high, and a wide range of cells can be infected. Their natural host cells are airway epithelium, so they are useful for therapy of lung cancers.

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Baculovirus-mediated transfer has several advantages. Baculoviral gene transfer can occur in replicating and nonreplicating cells, and can occur in renal cells, as well as in hepatocytes, neural cells, spleen, skin, and muscle. Baculovirus is non-replicating and nonpathogenic in mammalian cells. Humans lack pre-existing antibodies to recombinant baculovirus which could block infection. In addition, baculovirus is capable of incorporating and transducing very large DNA inserts.

Adeno-associated viruses (AAV) have also been employed as vectors for somatic gene therapy. AAV is a small, single-stranded (ss) DNA virus with a simple genomic organization (4-7 kb) that makes it an ideal substrate for genetic engineering. Two open reading frames encode a series of rep and cap polypeptides. Rep polypeptides (rep78, rep68, rep 62 and rep 40) are involved in replication, rescue and integration of the AAV genome. The cap proteins (VP1, VP2 and VP3) form the virion capsid. Flanking the rep and cap open reading frames at the 5' and 3' ends are 145 bp inverted terminal repeats (ITRs), the first 125 bp of which are capable of forming Y- or T-shaped duplex structures. Of importance for the development of AAV vectors, the entire rep and cap domains can be excised and replaced with a therapeutic or reporter transgene. See B.J. Carter, in Handbook of Parvoviruses, ed., P. Tijsser, CRC Press, pp. 155-168 (1990). It has been shown that the ITRs represent the minimal sequence required for replication, rescue, packaging, and integration of the AAV genome.

Adeno-associated viruses (AAV) have significant potential in gene therapy. The viral particles are very stable and recombinant AAVs (rAAV)have "drug-like" characteristics in that rAAV can be purified by pelleting or by CsCl gradient banding. They are heat stable and can be

- 29 -

lyophilized to a powder and rehydrated to full activity. Their DNA stably integrates into host chromosomes so expression is long-term. Their host range is broad and AAV causes no known disease so that the recombinant vectors are non-toxic.

Once introduced into a target cell, sequences of interest can be identified by conventional methods such as nucleic acid hybridization using probes comprising sequences that are homologous/complementary to the inserted gene sequences of the vector. In another approach, the sequence(s) may be identified by the presence or absence of a "marker" gene function (e.g, thymidine kinase activity, antibiotic resistance, and the like) caused by introduction of the expression vector into the target cell.

10 Formulations and Administration

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The compounds of the invention are formulated according to standard practice, such as prepared in a carrier vehicle. The term "pharmacologically acceptable carrier" means one or more organic or inorganic ingredients, natural or synthetic, with which the mutant proto-oncogene or mutant oncoprotein is combined to facilitate its application. A suitable carrier includes sterile saline although other aqueous and non-aqueous isotonic sterile solutions and sterile suspensions known to be pharmaceutically acceptable are known to those of ordinary skill in the art. In this regard, the term "carrier" encompasses liposomes and the HIV-1 tat protein (See Chen et al., Anal. Biochem. 227: 168-175, 1995) as well as any plasmid and viral expression vectors.

Any of the novel polypeptides of this invention may be used in the form of a pharmaceutically acceptable salt. Suitable acids and bases which are capable of forming salts with the polypeptides of the present invention are well known to those of skill in the art, and include inorganic and organic acids and bases.

A compound of the invention is administered to a subject in a therapeutically-effective amount, which means an amount of the compound which produces a medically desirable result or exerts an influence on the particular condition being treated. An effective amount of a compound of the invention is capable of ameliorating or delaying progression of the diseased, degenerative or damaged condition. The effective amount can be determined on an individual basis and will be based, in part, on consideration of the physical attributes of the subject,

- 30 -

symptoms to be treated and results sought. An effective amount can be determined by one of ordinary skill in the art employing such factors and using no more than routine experimentation.

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A liposome delivery system for a compound of the invention may be any of a variety of unilamellar vesicles, multilamellar vesicles, or stable plurilamellar vesicles, and may be prepared and administered according to methods well known to those of skill in the art, for example in accordance with the teachings of United States Patent 5,169,637, 4,762,915, 5,000,958 or 5,185,154. In addition, it may be desirable to express the novel polypeptides of this invention, as well as other selected polypeptides, as lipoproteins, in order to enhance their binding to liposomes. As an example, treatment of human acute renal failure with liposome-encapsulated KIM protein may be performed in vivo by introducing a KIM protein into cells in need of such treatment using liposomes. The liposomes can be delivered via catheter to the renal artery. The recombinant KIM protein is purified, for example, from CHO cells by immunoaffinity chromatography or any other convenient method, then mixed with liposomes and incorporated into them at high efficiency. The encapsulated protein may be tested in vitro for any effect on stimulating cell growth.

The compounds of the invention may be administered in any manner which is medically acceptable. This may include injections, by parenteral routes such as intravenous, intravascular, intraarterial, subcutaneous, intramuscular, intratumor, intraperitoneal, intraventricular, intraepidural, or others as well as oral, nasal, ophthalmic, rectal, or topical. Sustained release administration is also specifically included in the invention, by such means as depot injections or erodible implants. Localized delivery is particularly contemplated, by such means as delivery via a catheter to one or more arteries, such as the renal artery or a vessel supplying a localized tumor.

Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it will be obvious to one skilled in the art that certain changes and modifications may be practiced within the scope of the invention, as limited only by the scope of the appended claims.

WO 97/44460

- 31 -

SEQUENCE LISTING

(1) GENERAL INFORMATION:

(i) APPLICANT: Michele Sanicola-Nadel Joseph V. Bonventre Catherine A. Hession Takaharu Ichimura Henry Wei

Richard L. Cate

- (ii) TITLE OF INVENTION: MODULATORS OF TISSUE REGENERATION
- (iii) NUMBER OF SEQUENCES: 7
- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Biogen, Inc.
 - (B) STREET: 14 Cambridge Center
 - (C) CITY: Cambridge
 - (D) STATE: MA
 - (E) COUNTRY: USA
 - (F) ZIP: 02142
- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE: 23-MAY-1997
 - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 60/018,228
 - (B) FILING DATE: 24-MAY-1996
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Levine, Leslie M.
 - (B) REGISTRATION NUMBER: 35,245
 - (C) REFERENCE/DOCKET NUMBER: A010 PCT CIP
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (617) 679-2810
 - (B) TELEFAX: (617) 679-2838
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2566 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single

- 32 -

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 615..1535

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

GCGG	CCG	GT	CGAC	GTG	C TO	GTGAC	STAAF	A TAC	BATCI	AGGG	TCT	CTT	CAC	AGCA	CATTCI	60
CCAG	GAAC	SCC	GAGCI	AAACI	AT T	AGTG	TATI	TT)	ACCC2	AGGA	GGAJ	ATC:	rag (GTGT	AGAGAG	120
CTCI	racgo	EAT	CTAAC	GIT	rg g	ATCT	STACO	CAC	TGC:	TTT	TTAC	GTG:	rcr ·	TTAGI	ACATTI	180
CTCF	\GGAJ	AGA	TGTA	STCTO	T G	TCAC	CATGI	r GTC	GCT	TAAE	TCT	\GCT(CAG '	TCCA?	CTTAT	240
TYPTY		DAG	にかなご	יעכאי	ነር፤ ሞ	ኮ ሞክርረ	ממבי	יממ י	יראפי	באדמיו	TV*TV	TYCA(ברא (CARCI	AGTACA	300
			01110				aranc c	<i>.</i>	. دیمی	IAIG	1010		JCA V	GAMG	NOIACA	300
GTGI	rcca?	CT	TGAG	BACAI	KG C	TCAT	CTTT	r cci	ATTAC	BAGG	GCTC	GCC.	rtg (GCTT	AGATTO	360
TACC	GAG	AAC .	ATAC	rctc:	CA A	TGGC1	rgccc	: TC	\GTT	TCT	CTG	TTG	CTG '	TCTT	ATTTGT	420
GTC	\TGG(CCA	GAAG:	rcat;	AT GO	GATG	CTC1	TAT	3TGA(CAA	GGA	CCA	SAT :	AGAA	GAGTGT	480
ATT1	rece	GA ·	ACAGO	TTG(ec e:	TAACI	AGAGA	A GTO	CTG	rggg	ATTO	CATGO	CAG	TCAG	AADTAE	540
GACC	TGA	rca	GACA	BAGTO	T G	CTGAC	TGCC	C ACC	GCT1	AACC	AGAC	TGA	CTT (GTCA	CTGTCC	600
																
TICA	IGGT(:AA	CACC				Leu									650
				1	Val	GLII	Deu	5	val	FIIE	176	Ser	10	Deu	Deu	
Calca Calca	Calar	للملامل	CCA	GGC	TY-T	GTA	CAT	an Contr	ጥልጥ	CAA	CTT A	GTG.	AAG	ccc	CTC	698
			Pro													0.70
		15		,			20		-3-			25	-3-	,		
GTG	GGT	CAC	ССТ	GTC	ACA	ATT	CCA	TGT	ACT	TAC	TCA	ACA	CGT	GGA	GGA	746
			Pro													
	30					35					40					
ATC	ACA	ACG	ACA	TGT	TGG	GGC	CGG	GGG	CAA	TGC	CCA	TAT	TCT	AGT	TGT	794
	Thr	Thr	Thr	Сув	Trp	Gly	Arg	Gly	Gln	Сув	Pro	Tyr	Ser	Ser	Сув	
45					50					55					60	
CAA	AAT	ATA	CTT	ATT	TGG	ACC	AAT	GGA	TAC	CAA	GTC	ACC	TAT	CGG	AGC	842
Gln	Asn	Ile	Leu	Ile	Trp	Thr	Asn	Gly	Tyr	Gln	Val	Thr	Tyr	Arg	Ser	
				65					70					75		
AGC	GGT	CGA	TAC	AAC	ATA	AAG	GGG	CGT	ATT	TCA	GAA	GGA	GAC	GTA	TCC	890
Ser	Gly	Arg	Tyr	Asn	Ile	Lys	Gly	Arg	Ile	Ser	Glu	Gly	qaA	Val	Ser	
			80					85					90			

- 33 -

TTG	ACA	ATA	GAG	AAC	TCT	GTT	GAT	AGT	GAT	AGT	GGT	CIG	TAT	TGT	TGC	938
Leu	Thr	11e 95		Asn	Ser	Val	Asp 100	Ser	yab	Ser	Gly	Leu 105		Сув	Сув	
CGA	GTG	GAG	ATT	CCT	GGA	TGG	TTC	AAC	GAT	CAG	AAA	ATG	ACC	TTT	TCA	986
Arg	110	GIU	TTE	Pro	GIY	11p	Pne	Asn	Asp	Gln	. Lys 120		Thr	Phe	Ser	
TIG	GAA	GIT	AAA	CCA	GAA	ATT	CCC	ACA	AGT	CCT	CCA	ACA	AGA	CCC	ACA	1034
125	Giu	Val	гув	Pro	130	116	Pro	Thr	ser	Pro 135	Pro	Thr	Arg	Pro	Thr 140	
															-	
ACT	ACA	AGA	CCC	ACA	ACC	ACA	AGG	ccc	ACA	ACT	ATT	TCA	ACA	AGA	TCC	1082
Inr	Thr	arg	PTO	Thr 145	Thr	Thr	Arg	Pro	Thr 150	Thr	Ile	Ser	Thr		Ser	
														155		
ACA	CAT	GTA	CCA	ACA	TCA	ACC	AGA	GTC	TCC	ACC	TCT	ACT	CCA	ACA	CCA	1130
Thr	HIS	vaı	Pro 160	Thr	Ser	Thr	Arg	Val 165	Ser	Thr	Ser	Thr		Thr	Pro	
													170			
GAA	CAA	ACA	CAG	ACT	CAC	AAA	CCA	GAA	ATC	ACT	ACA	TTT	TAT	GCC	CAT	1178
GII	Gin	Thr 175	Gln	Thr	Ris	Lys	Pro 180	Glu	Ile	Thr	Thr		Tyr	Ala	His	•
							100					185				
GAG	ACA	ACT	GCT	GAG	GTG	ACA	GAA	ACT	CCA	TCA	TAT	ACT	CCT	GCA	GAC	1226
GIU	190	Thr	Ala	Glu	Val	Thr 195	Glu	Thr	Pro	Ser		Thr	Pro	Ala	Asp	
						193					200					
TGG	AAT	GGC	ACT	GTG	ACA	TCC	TCA	GAG	GAG	GCC	TGG	AAT	AAT	CAC	ACT	1274
11p 205	Asn	GTÅ	Thr	Val	Thr 210	Ser	Ser	Glu	Glu		Trp	Asn	Asn	His		
					210					215					220	
GTA	AGA	ATC	CCT	TTG	AGG	AAG	CCG	CAG	AGA	AAC	CCG	ACT	AAG	GGC	TTC	1322
vaı	Arg	IIe	Pro	Leu 225	Arg	Lys	Pro	Gln		Asn	Pro	Thr	rys	_	Phe	
									230					235		
TAT	GTT	GGC	ATG	TCC	GTT	GCA	GCC	CTG	CTG	CTG	CTG	CTG	CTT	GCG	AGC	1370
ıyr	vai	GIĀ	Met 240	Ser	Val	Ala	Ala	Leu 245	Leu	Leu	Leu	Leu		Ala	Ser	
													250			
ACC	GIG	GTT	GTC	ACC	AGG	TAC	ATC	ATT	ATA	AGA	AAG	DAA	ATG	GGC	TCT	1418
Inr	vai	va1 255	Val	Thr	Arg	Tyr	Ile 260	Ile	Ile	Arg	Lys		Met	Gly	Ser	
												265				
CTG	AGC	TTT	GTT	GCC	TTC	CAT	GTC	TCT	AAG	agt	AGA	GCT	TTG	CAG	AAC	1466
Leu	Ser 270	Phe	Val	Ala	Phe	His 275	Val	Ser	Lys	Ser		Ala	Leu	Gln	Asn	
	2.0					213					280					
GCA	GCG	ATT	GTG	CAT	CCC	CGA	GCT	GAA	GAC	AAC	ATC	TAC	ATT	ATT	GAA	1514
Ala 285	Ala	Ile	Val	His	Pro	Arg	Ala	Glu	Asp	Asn	Ile	Tyr	Ile	Ile	Glu	
-03					290					295					300	
GAT	AGA	TCT	CGA	GGT	GCA	GAA	TGAG	TCCC	AG A	GGCC	TTCT	G TG	GGGC	CTTC	<u>!</u>	1565
qaA	Arg	Ser	Arg	Gly .	Ala	Glu										

- 34 -

TG	CTGGGAT	TACAGAGATC	GTGACTGATT	TCACAGAGTA	AAATACCCAT	TCCAGCTCCT	1625
GG(BAGATTTT	GTGTTTTGGT	TCTTCCAGCT	GCAGTGGAGA	GGGTAACCCT	CTACCCTGTA	1685
TA?	rgcaaaac	TCGAGGTTAA	CATCATCCTA	ATTCTTGTAT	CAGCAACACC	TCAGTGTCTC	1745
CAC	TCACTGC	AGCGATTCTC	TCAAATGTGA	ACATTTTAGA	AGTTTGTGTT	TCCTTTTGTC	1805
CAT	rgtaatca	TTGGTAATAC	AAGAATTTTA	TCTTGTTTAT	TAAAACCATT	AATGAGAGGG	1865
GAJ	ATAGGAAT	TAAAAGCTGG	TGGGAAGGGC	CTCCTGAATT	TAGAAGCACT	TCATGATTGT	1925
GT	TATCTCT	TTTATTGTAA	TTTGAAATGT	TACTTCTATC	CTTCCCAAGG	GGCAAAATCA	1985
TGC	GAGCATG	GAGGTTTTAA	TIGCCCTCAT	AGATAAGTAG	AAGAAGAGAG	TCTAATGCCA	2045
CCI	AATAGAGG	TGGTTATGCT	TTCTCACAGC	TCTGGAAATA	TGATCATTTA	TTATGCAGTT	2105
GA7	PCTTAGGA	TGAGGATGGG	TTTCTTAGGA	GGAGAGGTTA	CCATGGTGAG	TGGACCAGGC	2165
ACJ	ACATCAGG	GGAAGAAAAC	AATGGATCAA	GGGATTGAGT	TCATTAGAGC	CATTTCCACT	2225
CCZ	CTTCTGT	CTTGATGCTC	AGTGTTCCTA	AACTCACCCA	CTGAGCTCTG	AATTAGGTGC	2285
AGC	GAGGAGA	CGTGCAGAAA	CGAAAGAGGA	AAGAAAGGAG	AGAGAGCAGG	ACACAGGCTT	2345
TC:	rgctgaga	GAAGTCCTAT	TGCAGGTGTG	ACAGTGTTTG	GGACTACCAC	GGGTTTCCTT	2405
CAC	BACTTCTA	AGTTTCTAAA	TCACTATCAT	GTGATCATAT	TTATTTTTAA	AATTATTTCA	2465
GAJ	AGACACC	ACATTTTCAA	TAATAAATCA	GTTTGTCACA	ATTAATAAAA	TATTITGTTT	2525
GCT	TAAGAAGT	АААААААА	AAAAAAAGTC	GACGCGGCCG	C		2566

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2084 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 145..1065
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

- 35 -

CCA	GGAA	GCC	GAGC	AAAC	AT T	AGTG	CTAT	T T	TACCC	AGGA	A GGZ	YEAA	TAG	GTG:	ragaga	rg	120
CTC	TACG	GAT	CTAA	GGTC	AA C			GTT Val									171
GGC Gly 10	Leu	Leu	CTG Leu	CTT Leu	CTT Leu 15	Pro	GGC	TCT Ser	GTA Val	GAT Asp 20	Sex	TAT	GAZ Glu	GTZ Val	GTG Val 25		219
AAG Lys	GGG	GTG Val	GTG Val	GGT Gly 30	CAC His	Pro	GTC Val	ACA Thr	ATT Ile 35	Pro	TGT Cys	Thr	TAC	Ser 40	ACA Thr	·	267
CGT Arg	GGA Gly	GGA Gly	Ile 45	Thr	ACG Thr	ACA	TGT Cys	TGG Trp 50	Gly	CGG Arg	GGG Gly	Gln	TGC Cys 55	Pro	TAT Tyr		315
TCT Ser	AGT Ser	TGT Cys 60	Gln	AAT Asn	ATA Ile	CTT	ATT Ile 65	Trp	ACC	AAT Asn	GGA Gly	TAC Tyr 70	Gln	GTC Val	ACC		363
TAT Tyr	CGG Arg 75	AGC Ser	AGC Ser	GGT Gly	CGA Arg	TAC Tyr 80	AAC	ATA Ile	AAG Lys	GGG	CGT Arg 85	Ile	TCA Ser	GAA Glu	GGA Gly		411
GAC Asp 90	GTA Val	TCC	TTG Leu	ACA Thr	ATA Ile 95	GAG Glu	AAC	TCT	GTT Val	GAT Asp 100	AGT Ser	GAT Asp	AGT Ser	GGT Gly	Leu 105		459
TAT Tyr	TGT Cys	TGC Cys	CGA Arg	GTG Val 110	GAG Glu	ATT Ile	CCT Pro	GGA Gly	TGG Trp 115	TTC Phe	AAC Asn	GAT Asp	CAG Gln	AAA Lys 120	ATG Met	!	507
ACC Thr	TTT Phe	TCA Ser	TTG Leu 125	GAA Glu	GTT Val	AAA Lys	CCA Pro	GAA Glu 130	ATT Ile	CCC	ACA Thr	AGT Ser	CCT Pro 135	CCA	ACA Thr	!	555
AGA Arg	Pro	ACA Thr 140	ACT Thr	ACA Thr	AGA Arg	CCC Pro	ACA Thr 145	ACC Thr	ACA Thr	AGG Arg	CCC	ACA Thr 150	ACT Thr	ATT Ile	TCA Ser	•	503
ACA Thr	AGA Arg 155	TCC Ser	ACA Thr	CAT His	GTA Val	CCA Pro 160	ACA Thr	TCA Ser	ACC Thr	AGA Arg	GTC Val 165	TCC Ser	ACC Thr	TCT Ser	ACT Thr	•	551
CCA Pro 170	ACA Thr	CCA Pro	GAA Glu	CAA Gln	ACA Thr 175	CAG Gln	ACT Thr	CAC His	AAA Lys	CCA Pro 180	GAA Glu	ATC Ile	ACT Thr	ACA Thr	TTT Phe 185	€	599
TAT Tyr	GCC Ala	CAT His	GAG Glu	ACA Thr 190	ACT Thr	GCT Ala	GAG Glu	GTG Val	ACA Thr 195	GAA Glu	ACT Thr	CCA Pro	TCA Ser	TAT Tyr 200	ACT Thr	7	147

- 36 -

														TGG Trp	AAT Asn	795
														CCG Pro		843
														CTG Leu		891
														AAG Lys		939
														AGA Arg 280		987
														ATC Ile		1035
			GAT Asp							TGAG	FTCC	CAG A	AGGC	CTTC	rg	1085
TGG	GCC	TTC :	rgcc	rgggj	AT T	ACAGI	AGATO	C GTC	SACTO	TTAE	TCAC	CAGA	STA Z	AAATI	ACCCAT	1145
TCC	AGCTO	CT (GGGA	SATT	rt g	GTT	rrgg	r TCT	TCC	AGCT	GCAC	TGG	AGA (GGT1	AACCCI	1205
CTA	CCTC	TA :	TATG	CAAA	AC TO	CGAG	STTAI	A CAT	CATO	CTA	ATTO	TTG	TAT	CAGC	AACACC	1265
TCA	TGT	TC (CACTO	CACTY	SC A	CGA:	rrcr	TC	YAAF	FTGA	ACA?	TTT1	AGA	AGTT	IGTGTT	1325
TCC	CTTT	TC (CATG	raat(CA T	rggty	AATA	C AAC	BAATT	PTTA	TCT	GTT	TAT	TAAAI	ACCATI	1385
AAT	BAGA	egg (Gaati	AGGAJ	AT T	AAAA	CTG	3 TGC	3GAA (3GGC	CTC	TGA	ATT '	TAGAI	AGCACI	1445
TCA:	GAT.	rgt (GTTT!	ATCTO	T T	TAT	IGTA	A TT	(AAD1	ATGT	TACT	TCT	ATC (CTTC	CCAAGG	1505
GGCI	AAAA?	rca :	TGGG	AGCA?	rg g	AGGT	TTTAJ	A TTO	ECCC!	CAT	AGA	raag:	Mag :	AAGA	AGAGAG	1565
TCT	AATG	CCA (CCAA!	PAGA	G T	GTT	ATGC:	r TT	CTCA	CAGC	TCT	GAA	ATA '	TGAT	CATTTA	1625
TTA:	rgcad	err (GATC.	TAG	JA T	BAGG	ATGG	3 TT	CTT	AGGA	GGA	GAGG"	rta	CCAT	GTGAG	1685
TGG	ACCA	GC 2	ACACI	ATCA	GG G	BAAG	AAAA	C AA	rgga:	ICAA	GGGJ	ATTG/	AGT '	TCAT	ragago	1745
CAT	rtcc.	ACT (CCAC	TCI	T C	rtga:	IGCT	C AG	IGTT	CCTA	AAC	CAC	CCA	CTGA	CTCT	1805
AAT	ragg:	rgc i	AGGG	AGGA	BA C	GTGC:	AGAA	A CG	NAAG	AGGA	AAG	AAA G(GAG .	AGAG	AGCAGO	1865
ACA	CAGG	CTT '	TCTG	CTGA	GA G	aagty	CCTA:	r TG	CAGG'	rgrg	ACA	TGT:	TTG	GGAC	TACCAC	1925

- 37 -

GGGTTTCCTT	CAGACTTCTA	AGTTTCTAAA	TCACTATCAT	GTGATCATAT	TTATTTTAA	1985
AATTATTTCA	GAAAGACACC	ACATTTTCAA	ТААТАААТСА	GTTTGTCACA	ATTAATAAA	2045
TATTTTGTTT	GCTAAGAAGT	AAAAAGTCGA	ceceeccec			2084
(2) INFORME	ATION POR SE	O ID NO.3.				

RMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 307 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

Met Val Gln Leu Gln Val Phe Ile Ser Gly Leu Leu Leu Leu Pro

Gly Ser Val Asp Ser Tyr Glu Val Val Lys Gly Val Val Gly His Pro 25

Val Thr Ile Pro Cys Thr Tyr Ser Thr Arg Gly Gly Ile Thr Thr 40

Cys Trp Gly Arg Gly Gln Cys Pro Tyr Ser Ser Cys Gln Asn Ile Leu

Ile Trp Thr Asn Gly Tyr Gln Val Thr Tyr Arg Ser Ser Gly Arg Tyr

Asn Ile Lys Gly Arg Ile Ser Glu Gly Asp Val Ser Leu Thr Ile Glu

Asn Ser Val Asp Ser Asp Ser Gly Leu Tyr Cys Cys Arg Val Glu Ile

Pro Gly Trp Phe Asn Asp Gln Lys Met Thr Phe Ser Leu Glu Val Lys 120

Pro Glu Ile Pro Thr Ser Pro Pro Thr Arg Pro Thr Thr Thr Arg Pro 135

Thr Thr Thr Arg Pro Thr Thr Ile Ser Thr Arg Ser Thr His Val Pro 150

Thr Ser Thr Arg Val Ser Thr Ser Thr Pro Thr Pro Glu Gln Thr Gln 170

Thr His Lys Pro Glu Ile Thr Thr Phe Tyr Ala His Glu Thr Thr Ala

PCT/US97/09303 WO 97/44460

	20	
•	20	-

									- 31	S -			•			
Glu	Val	Thr 195	Glu	Thr	Pro	Ser	Tyr 200	Thr	Pro	Ala	Авр	Trp 205	Asn	Gly	Thr	
Val	Thr 210	Ser	Ser	Glu	Glu	Ala 215	Trp	Asn	Asn	His	Thr 220	Val	Arg	Ile	Pro	
Leu 225	Arg	Lys	Pro	Gln	Arg 230	Asn	Pro	Thr	Lys	Gly 235	Phe	Tyr	Val	Gly	Met 240	
Ser	Val	Ala	Ala	Leu 245	Leu	Leu	Leu	Leu	Leu 250	Ala	Ser	Thr	Val	Val 255	Val	
Thr	Arg	Tyr	1le 260	Ile	Ile	Arg	Lys	Lys 265	Met	Gly	Ser	Leu	Ser 270	Phe	Val	
Ala	Phe	His 275	Val	Ser	Lys	Ser	Arg 280	Ala	Leu	Gln	Asn	Ala 285	Ala	Ile	Val	
His	Pro 290	Arg	Ala	Glu	Asp	Asn 295	Ile	Tyr	Ile	Ile	Glu 300	Asp	Arg	Ser	Arg	
Gly 305	Ala	Glu														
(2)	INFO	ORMA?	CION	FOR	SEQ	ID 1	10:4 :	:								
		() () (1)	A) Li 3) T: C) S: O) T(engti YPE : I'RANI OPOLA	nucl DEDNI OGY:	303 l leic 3SS: line	ase acid sing ar	pai:	ra							
	(11,) MOI	LECUI	LE T	KAR:	CDM	•									
	(ix		A) NI	ame/i	KEY: ION:		18:	22								
	(xi) SE()OBN	CR DI	BSCR:	[PTI	ON:	SEQ :	ID N	0:4:						
GCG	GCCG	CGT (CGAC.	rcgc:	AG G	AGGC	CGGC	A CT	CTGA	CTCC	TGG	rgga'	rgg (3ACT/	AGGGA	G 60
TCA	gagt	CAA (GCCC.	TGAC:	rg ge	CTGA	GGC	G GG	CGCT	CCGA	GTC		ATG (Met (115
		GGG														163
	Ala	GCC Ala			-											211

- 39 -

GAT Asp	CAC His	Met	AGG Arg	GAG Glu 40	Asn	AAC Asn	CAA Gln	Leu	CGT Arg 45	Gly	Trp	Ser	TCA Ser	GAT Asy 50	GAA Glu	259
AAT Asn	GAA	TGG Trp	GAT Asp 55	Glu	CAG Gln	CTG Leu	TAT	CCA Pro 60	Val	TGG	AGG	AGG Arg	GGA Gly 65	Glu	GGC Gly	301
AGA Arg	TGG	AAG Lys	Asp	TCC Ser	TGG Trp	GAA Glu	GGA Gly 75	Gly	CGT Arg	GTG Val	CAG Gln	GCA Ala 80	Ala	CTA Leu	ACC Thr	355
AGT Ser	GAT Asp 85	Ser	CCG Pro	GCC	TTG Leu	GTG Val 90	GGT Gly	TCC	AAT Asn	ATC Ile	ACC Thr 95	Phe	GTA Val	GTG Val	AAC	403
CTG Leu 100	GTG Val	TTC	Pro	AGA Arg	TGC Cys 105	CAG Gln	AAG Lys	GAA Glu	GAT Asp	GCC Ala 110	AAC Asn	GGC Gly	AAT Asn	ATC Ile	GTC Val 115	451
TAT Tyr	GAG Glu	AGG	AAC Asn	TGC Cys 120	AGA Arg	AGT Ser	GAT Asp	TTG Leu	GAG Glu 125	CTG Leu	GCT Ala	TCT Ser	GAC Asp	CCG Pro 130	TAT Tyr	499
GTC Val	TAC Tyr	AAC Asn	TGG Trp 135	ACC Thr	ACA Thr	GGG Gly	GCA Ala	GAC Asp 140	GAT Asp	GAG Glu	GAC Asp	TGG	GAA Glu 145	GAC Asp	AGC Ser	547
ACC Thr	AGC Ser	CAA Gln 150	GGC	CAG Gln	CAC His	CTC Leu	AGG Arg 155	TTC Phe	CCC Pro	GAC Asp	GGG Gly	AAG Lys 160	CCC Pro	TTC Phe	CCT Pro	595
CGC Arg	CCC Pro 165	CAC His	GGA Gly	CGG Arg	AAG Lys	AAA Lys 170	TGG Trp	AAC Asn	TTC Phe	GTC Val	TAC Tyr 175	GTC Val	TTC Phe	CAC His	ACA Thr	643
CTT Leu 180	GGT Gly	CAG Gln	TAT Tyr	TTT Phe	CAA Gln 185	AAG Lys	CTG Leu	ggt Gly	CGG Arg	ТСТ Сув 190	TCA Ser	GCA Ala	CGA Arg	GTT Val	TCT Ser 195	691
ATA Ile	AAC Asn	ACA Thr	GTC Val	AAC Asn 200	TTG Leu	ACA Thr	GTT Val	GGC Gly	CCT Pro 205	CAG Gln	GTC Val	ATG Met	GAA Glu	GTG Val 210	ATT Ile	739
GTC Val	TTT Phe	CGA Arg	AGA Arg 215	CAC His	GGC	CGG Arg	GCA Ala	TAC Tyr 220	ATT Ile	CCC Pro	ATC Ile	Ser	AAA Lys 225	GTG Val	aaa Lys	787
gac Asp	GTG Val	TAT Tyr 230	GTG Val	ATA Ile	ACA Thr	Asp	CAG Gln 235	ATC Ile	CCT Pro	ATA Ile	Phe	GTG Val 240	ACC Thr	ATG Met	TAC Tyr	835
GIN	AAG Lys 245	AAT Asn	GAC Asp	CGG Arg	Asn	TCG Ser	TCT Ser	GAT Asp	GAA . Glu	Thr	TTC Phe 255	CTC . Leu .	AGA Arg	GAC Asp	CTC Leu	883

- 40 -

CCC	ATT	TTC	TTC	GAT	GTC	CTC	ATT	CAC	GAT	CCC	AGT	CAT	TTC	CTC	AAC	931
	Ile	Phe	Phe	Авр	Val	Leu	Ile	His	Asp	Pro	Ser	His	Phe	Leu	Asn	
260					265					270					275	
			ATT													979
ıyr	ser	Ala	Ile		Tyr	Lys	Trp	Asn		Gly	Asp	asa	Thr	_	Leu	
				280					285					290		
defete	OTC.	TO CO	AAC		CD C	200			~~~							
			Asn													1027
FIIC	Val	Ser	295	MBII	UTR	IIIK	Den	300	HIS	Inr	туг	vaı	305	Asn	GIA	
								300					303			
ACC	TTC	AAC	TIT	AAC	CTC	ACC	GTG	CAA	ACT	GCA	GTG	CCG	GGA	CCA	TCC	1075
			Phe													1075
		310					315					320	 ,		Cyb	
CCC	TCA	CCC	ACA	CCT	TCG	CCT	TCT	TCT	TCG	ACT	TCT	CCT	TCG	CCT	GCA	1123
Pro	Ser	Pro	Thr	Pro	Ser	Pro	Ser	Ser	Ser	Thr	Ser	Pro	Ser	Pro	Ala	
	325					330					335					
			TCA													1171
	Ser	Pro	Ser	Pro		Leu	Ser	Thr	Pro		Pro	Ser	Leu	Met	Pro	
340					345					350					355	
N/T	ccc	CAC		maa	3.000	~~~	~~~		~~~							
			AAA Lys													1219
****	GLY	1110	Lys	360	MCL	GIU	Deu	SEL	365	116	ser	ASII	GIU		Сув	
				500					303					370		
CGA	ATA	AAC	AGA	TAT	GGT	TAC	TTC	AGA	GCC	ACC	ATC	ACA	ATT	ATD.	GAT	1267
			Arg													1207
_			375	•	•	•		380					385			
GGA	ATC	CTA	GAA	GTC	AAC	ATC	ATC	CAG	GTA	GCA	GAT	GTC	CCA	ATC	CCC	1315
Gly	Ile	Leu	Glu	Val	Asn	Ile	Ile	Gln	Val	Ala	Asp	Val	Pro	Ile	Pro	
		390					395					400				
			CCT													1363
THE	405	GIN	Pro	Авр	Asn		Leu	Met	Asp	Phe		Val	Thr	Сув	Lys	
	403					410					415					
GGG	GCC	ארידי	ccc	ACC.	GAA	acc	THE	»CC	አጥሮ	አጥሮ	TV-T	CAC	ccc	200	TCC	2422
			Pro													1411
420					425	-	Cyb		110	430	BCI	veh	PIO	1111	435	
															433	
CAG	ATC	GCC	CAG	AAC	AGG	GTG	TGC	AGC	CCG	GTG	GCT	GTG	GAT	GAG	CTG	1459
			Gln													
				440			-		445			•	•	450		
			TCC													1507
Сув	Leu	Leu	Ser	Val	Arg	Arg	Ala	Phe	Asn	Gly	Ser	Gly	Thr	Tyr	Сув	
			455					460					465			•
			ACT													1555
vaı	ABN		Thr	Leu	GIA	Asp		Ala	Ser	Leu	Ala		Thr	Ser	Ala	
		470					475					480				

- 41 -

CTG Leu	ATC Ile 485	TCT Ser	ATC Ile	CCT Pro	GGC Gly	AAA Lys 490	GAC Asp	CTA Leu	GGC	TCC Ser	CCT Pro 495	CTG Leu	AGA Arg	ACA Thr	GTG Val		1603
AAT Asn 500	Gly	GTC Val	CTG Leu	ATC Ile	TCC Ser 505	ATT Ile	GGC Gly	TGC Cys	CTG Leu	GCC Ala 510	ATG Met	TTT Phe	GTC Val	ACC Thr	ATG Met 515		1651
GTT Val	ACC Thr	ATC Ile	TTG Leu	CTG Leu 520	TAC Tyr	AAA Lys	AAA Lyb	CAC His	AAG Lys 525	ACG Thr	TAC Tyr	AAG Lys	CCA Pro	ATA Ile 530	GGA Gly		1699
AAC Asn	TGC Cys	ACC Thr	AGG Arg 535	AAC Asn	GTG Val	GTC Val	AAG Lys	GGC Gly 540	AAA Lys	GGC Gly	CTG Leu	AGT Ser	GTT Val 545	TTT Phe	CTC Leu		1747
AGC Ser	CAT His	GCA Ala 550	AAA Lys	GCC Ala	CCG Pro	TTC Phe	TCC Ser 555	CGA Arg	GGA Gly	GAC Asp	CGG Arg	GAG Glu 560	AAG Lys	GAT Asp	CCA Pro		1795
						TGG Trp 570		CTC Leu	TAAG	TCTT	CA C	TCTC	ACTI	rc _.			1842
TGAC	TGGG	AA C	CCAC	TCTI	C TO	TGC	TGT	TGI	GAGC	TGT	GCAG	AAGT	AC A	TGAC	TGGTA		1902
GCTG	TTGT	TT 1	CTAC	GGAT	T AT	TGTA	CAAA	GTA	TATO	ATG	GTTI	'AGGG	IAG C	GTAG	TTAAT		1962
TGGC	ATT1	TA G	TGA	AGGGA	T GG	GAAG	ACAG	TAT	TTCT	TCA	CATO	TGTA	TT G	TGGT	TTTTA		2022
TACI	GTTA	AT A	\GGG1	regec	A CA	TTGT	GICI	GAA	œœ	GAG	GGGG	AGGT	CA C	TGCI	ACTTA		2082
AGG'I	CCTA	LGG 1	TAAC	TGGG	ia ga	GGAI	GCCC	CAG	GCTC	CTT	AGAT	TTCI	'AC A	CAAG	ATGTG		2142
CCTG	IAACC	CA G	CTAG	TCCI	G AC	CTAA	AGGC	CAI	GCTT	CAT	CAAC	TCTA	TC I	CAGO	TCATT		2202
GAAC	ATAC	CT G	BAGCA	CCTG	A TO	TAAD	TATA	ATG	GAAC	CAA	GCTI	GTTG	TA I	GGTG	TGTGT		2262
GTGI	'ACAT	'AA G	ATAC	TCAT	T A	AAAG	ACAG	TCT	ATTA	AAA	A					;	2303

(2) INFORMATION FOR SEQ ID NO:5:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 572 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Met Glu Ser Leu Cys Gly Val Leu Val Phe Leu Leu Leu Ala Ala Gly

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PCT/US97/09303 WO 97/44460

- 42 -

Leu Pro Leu Gln Ala Ala Lys Arg Phe Arg Asp Val Leu Gly His Glu 25

Gln Tyr Pro Asp His Met Arg Glu Asn Asn Gln Leu Arg Gly Trp Ser

Ser Asp Glu Asn Glu Trp Asp Glu Gln Leu Tyr Pro Val Trp Arg Arg

Gly Glu Gly Arg Trp Lys Asp Ser Trp Glu Gly Gly Arg Val Gln Ala

Ala Leu Thr Ser Asp Ser Pro Ala Leu Val Gly Ser Asn Ile Thr Phe

Val Val Asn Leu Val Phe Pro Arg Cys Gln Lys Glu Asp Ala Asn Gly

Asn Ile Val Tyr Glu Arg Asn Cys Arg Ser Asp Leu Glu Leu Ala Ser

Asp Pro Tyr Val Tyr Asn Trp Thr Thr Gly Ala Asp Asp Glu Asp Trp 135

Glu Asp Ser Thr Ser Gln Gly Gln His Leu Arg Phe Pro Asp Gly Lys

Pro Phe Pro Arg Pro His Gly Arg Lys Lys Trp Asn Phe Val Tyr Val

Phe His Thr Leu Gly Gln Tyr Phe Gln Lys Leu Gly Arg Cys Ser Ala

Arg Val Ser Ile Asn Thr Val Asn Leu Thr Val Gly Pro Gln Val Met 200

Glu Val Ile Val Phe Arg Arg His Gly Arg Ala Tyr Ile Pro Ile Ser

Lys Val Lys Asp Val Tyr Val Ile Thr Asp Gln Ile Pro Ile Phe Val

Thr Met Tyr Gln Lys Asn Asp Arg Asn Ser Ser Asp Glu Thr Phe Leu 250

Arg Asp Leu Pro Ile Phe Phe Asp Val Leu Ile His Asp Pro Ser His 260 265

Phe Leu Asn Tyr Ser Ala Ile Ser Tyr Lys Trp Asn Phe Gly Asp Asn

Thr Gly Leu Phe Val Ser Asn Asn His Thr Leu Asn His Thr Tyr Val 290 295

- 43 -

Leu	Asn	Gly	Thr	Phe	Asn	Phe	Asn	Leu	Thr	Val	Gln	Thr	Ala	Val	Pro
305					310					315					320

- Gly Pro Cys Pro Ser Pro Thr Pro Ser Pro Ser Ser Ser Thr Ser Pro 325 330 335
- Ser Pro Ala Ser Ser Pro Ser Pro Thr Leu Ser Thr Pro Ser Pro Ser 340 345 350
- Leu Met Pro Thr Gly His Lys Ser Met Glu Leu Ser Asp Ile Ser Asn 355 360 365
- Glu Asn Cys Arg Ile Asn Arg Tyr Gly Tyr Phe Arg Ala Thr Ile Thr 370 375 380
- Ile Val Asp Gly Ile Leu Glu Val Asn Ile Ile Gln Val Ala Asp Val 385 390 395 400
- Pro Ile Pro Thr Pro Gln Pro Asp Asn Ser Leu Met Asp Phe Ile Val 405 410 415
- Thr Cys Lys Gly Ala Thr Pro Thr Glu Ala Cys Thr Ile Ile Ser Asp 420 425 430
- Pro Thr Cys Gln Ile Ala Gln Asn Arg Val Cys Ser Pro Val Ala Val 435 440 445
- Asp Glu Leu Cys Leu Leu Ser Val Arg Arg Ala Phe Asn Gly Ser Gly 450 455 460
- Thr Tyr Cys Val Asn Phe Thr Leu Gly Asp Asp Ala Ser Leu Ala Leu 465 470 475 480
- Thr Ser Ala Leu Ile Ser. Ile Pro Gly Lys Asp Leu Gly Ser Pro Leu 485 490 495
- Arg Thr Val Asn Gly Val Leu Ile Ser Ile Gly Cys Leu Ala Met Phe 500 505 510
- Val Thr Met Val Thr Ile Leu Leu Tyr Lys Lys His Lys Thr Tyr Lys 515 520 525
- Pro Ile Gly Asn Cys Thr Arg Asn Val Val Lys Gly Lys Gly Leu Ser 530 540
- Val Phe Leu Ser His Ala Lys Ala Pro Phe Ser Arg Gly Asp Arg Glu 545 550 555 560
- Lys Asp Pro Leu Leu Gln Asp Lys Pro Trp Met Leu 565 570
- (2) INFORMATION FOR SEQ ID NO:6:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1795 base pairs

- 44 -

(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(ix) FRATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 278..1279

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

GCGG	CCGC	GT (CGACG	AAGO	T GO	GAAC	TCAC	GGG	CTGI	TTC	TGTG	GGCZ	AGC 1	TTC	CTGTC	60
CTTI	GGAA	GG (CACAG	AGC	C TO	AGCI	GCAG	GG2	ACTA	ACA	GAGO	TCT	AA G	ccca	TATAT	120
GTGG	TCTI	CT (CTCAT	TTC	A GO	CAGAG	CAGO	CTC	CATAI	'GAA	TCA	CCA	CT G	GGT	BAAAAG	180
ATAA	GTTG	CA 2	ATCTO	GAGA'	T T	AGAC	TTG	TC	GATA	CCA	TCTC	GTG	AG C	GTAC	CAACC	240
AGCC	TGTC	TG (CTCA1	TTT	C T	CAGG	ectg <i>i</i>	A TCC	CATA		: His				l Val	295
ATC	TTA	AGC	CTC	ATC	CTA	CAT	CTG	GCA	GAT	TCT	GTA	GCT	GGT	TCT	GTA	343
Ile	Leu	Ser	Leu 10	Ile	Leu	His	Leu	Ala 15	Авр	Ser	Val	Ala	Gly 20	Ser	Val	
			GGA													391
Lys	Val	Gly 25	Gly	Glu	Ala	Gly	Pro 30	Ser	Val	Thr	Leu	Pro 35	Сув	His	Tyr	
agt	GGA	GCT	GTC	ACA	TCA	ATG	TGC	TGG	AAT	AGA	GGC	TCA	TGT	TCT	CTA	439
Ser	Gly 40	Ala	Val	Thr	Ser	Met 45	Сув	Trp	Asn	Arg	Gly 50	Ser	Сув	Ser	Leu	
TTC	ACA	TGC	CAA	AAT	GGC	ATT	GTC	TGG	ACC	AAT	GGA	ACC	CAC	GTC	ACC	487
Phe 55	Thr	Сув	Gln	Asn	Gly 60	Ile	Val	Trp	Thr	Asn 65	Gly	Thr	His	Val	Thr 70	
TAT	CGG	AAG	GAC	ACA	CGC	TAT	AAG	CTA	TTG	GGG	GAC	CTT	TCA	AGA	AGG	535
Tyr	Arg	ГÀа	Asp	Thr 75	Arg	Tyr	Lys	Leu	Leu 80	Gly	Asp	Leu	Ser	Arg 85	Arg	
GAT	GTC	TCT	TTG	ACC	ATA	GAA	AAT	ACA	GCT	GTG	TCT	GAC	AGT	GGC	GTA	583
Авр	Val	Ser	Leu 90	Thr	Ile	Glu	Asn	Thr 95	Ala	Val	Ser	Двр	Ser 100	Gly	Val	
TAT	TGT	TGC	CGT	GTT	GAG	CAC	CGT	GGG	TGG	TTC	AAT	GAC	ATG	AAA	ATC	631
Tyr	Сув	Cys 105	Arg	Val	Glu	His	Arg 110	_	Trp	Phe	Asn	Авр 115	Met	Lys	Ile	

- 45 -

ACC Thr	GTA Val 120	Ser	TTG Leu	GAG Glu	ATT	GTG Val 125	Pro	CCC	AAG Lys	GTC Val	ACG Thr	Thr	ACT Thr	CCA Pro	ATT	679
GTC Val 135	Thr	ACT	GTT Val	CCA Pro	ACC Thr 140	Val	ACG Thr	ACT Thr	GTT Val	CGA Arg 145	Thr	AGC Ser	ACC Thr	ACT	GTT Val 150	727
CCA Pro	ACG Thr	ACA Thr	ACG Thr	ACT Thr 155	GTT Val	CCA Pro	ACG Thr	ACA Thr	ACT Thr 160	GTT Val	CCA Pro	ACA Thr	ACA Thr	ATG Met 165	AGC Ser	775
ATT	CCA Pro	ACG Thr	ACA Thr 170	ACG Thr	ACT Thr	GTT Val	CCG Pro	ACG Thr 175	ACA Thr	ATG Met	ACT Thr	GTT Val	TCA Ser 180	ACG Thr	ACA Thr	823
ACG Thr	AGC Ser	GTT Val 185	CCA Pro	ACG Thr	ACA Thr	ACG Thr	AGC Ser 190	ATT Ile	CCA Pro	ACA Thr	ACA Thr	ACA Thr 195	AGT Ser	GTT Val	CCA Pro	871
Val	Thr 200	Thr	Thr	Val	Ser	Thr 205	Phe	Val	CCT Pro	Pro	Met 210	Pro	Leu	Pro	Arg	919
Gln 215	Asn	His	Glu	Pro	Val 220	Ala	Thr	Ser	CCA Pro	Ser 225	Ser	Pro	Gln	Pro	Ala 230	967
Glu	Thr	His	Pro	Thr 235	Thr	Leu	Gln	Gly	GCA Ala 240	Ile	Arg	Arg	Glu	Pro 245	Thr	1015
Ser	Ser	Pro	Leu 250	Tyr	Ser	Tyr	Thr	Thr 255	Asp Asp	Gly	Asn	Asp	Thr 260	Val	Thr	1063
Glu	Ser	Ser 265	qaA	Gly	Leu	Trp	Asn 270	Asn	AAT Asn	Gln	Thr	Gln 275	Leu	Phe	Leu	1111
Glu	His 280	Ser	Leu	Leu	Thr	Ala 285	Asn	Thr	ACT Thr	Lys	Gly 290	Ile	Tyr	Ala	Gly	1159
Val 295	Сув	Ile	Ser	Val	Leu 300	Val	Leu	Leu	GCT Ala	Leu 305	Leu	Gly	Val	Ile	Ile 310	1207
Ala	Lys	Lys	Tyr	Phe 315	Phe	Lys	Lys	Glu	GTT Val 320	Gln	Gln	Leu	Arg	Pro 325	His	1255
AAA Lys	TCC Ser	TGT Cys	ATA Ile 330	CAT His	CAA Gln	AGA Arg	GAA Glu	TAGT	CCCT	GG A	AACA	TAGC	A AA	TGAA	CTTC	1309

- 46 -

**************************************	ATCACACCTC	TCCAGAAGAG	CCCD DTCTCT	CHERNANACC	AGCAAATCCA	1260
INICIIOGCC	ATCACAGE 1G	1 CCROPINGING	GGGAATCIGI	CIIAMANCC	AGCAAATCCA	1309
ACGTGAGACT	TCATTTGGAA	GCATTGTATG	ATTATCTCTT	GTTTCTATGT	TATACTTCCA	1429
AATGTTGCAT	TTCCTATGTT	TTCCAAAGGT	TTCAAATCGT	GGGTTTTTAT	TTCCTCCGTG	1489
GGGAAACAAA	GTGAGTCTAA	CTCACAGGTT	TAGCTGTTTT	CTCATAACTC	TGGAAATGTG	1549
ATGCATTAAG	TACTGGATCT	CTGAATTGGG	GTAGCTGTTT	TACCAGTTAA	AGAGCCTACA	1609
atagtatgga	ACACATAGAC	ACCAGGGGAA	GAAAATCATT	TGCCAGGTGA	TTTAACATAT	1669
TTATGCAATT	TTTTTTTTT	TTTTTGAGAT	GGAGCTTTGC	TCTTGTTGCC	CAGGCTGGAG	1729
TGCGATGGTG	AAATCTCGGC	TCACTGTAAC	CTCCACCTTC	CGGGTTCAAG	CAATTCTCCC	1789
GTCGAC						1795

(2) INFORMATION FOR SEQ ID NO:7:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 334 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Met His Pro Gln Val Val Ile Leu Ser Leu Ile Leu His Leu Ala Asp 1 5 10 15

Ser Val Ala Gly Ser Val Lys Val Gly Glu Ala Gly Pro Ser Val 20 25 30

Thr Leu Pro Cys His Tyr Ser Gly Ala Val Thr Ser Met Cys Trp Asn 35 40 45

Arg Gly Ser Cys Ser Leu Phe Thr Cys Gln Asn Gly Ile Val Trp Thr 50 55 60

Asn Gly Thr His Val Thr Tyr Arg Lys Asp Thr Arg Tyr Lys Leu Leu 65 70 75 80

Gly Asp Leu Ser Arg Arg Asp Val Ser Leu Thr Ile Glu Asn Thr Ala 85 90 95

Val Ser Asp Ser Gly Val Tyr Cys Cys Arg Val Glu His Arg Gly Trp 100 105 110

Phe Asn Asp Met Lys Ile Thr Val Ser Leu Glu Ile Val Pro Pro Lys 115 120 125

- 47 -

Val	Thr	Thr	Thr	Pro	Ile	Val	Thr	Thr	Val	Pro	Thr	Val	Thr	Thr	Val
	130					135					140				

- Arg Thr Ser Thr Thr Val Pro Thr Thr Thr Thr Val Pro Thr Thr Thr 145 150 155 160
- Val Pro Thr Thr Met Ser Ile Pro Thr Thr Thr Thr Val Pro Thr Thr 165 170 175
- Met Thr Val Ser Thr Thr Thr Ser Val Pro Thr Thr Thr Ser Ile Pro 180 185 190
- Thr Thr Thr Ser Val Pro Val Thr Thr Thr Val Ser Thr Phe Val Pro 195 200 205
- Pro Met Pro Leu Pro Arg Gln Asn His Glu Pro Val Ala Thr Ser Pro 210 215 220
- Ser Ser Pro Gln Pro Ala Glu Thr His Pro Thr Thr Leu Gln Gly Ala 225 230 235 240
- Ile Arg Arg Glu Pro Thr Ser Ser Pro Leu Tyr Ser Tyr Thr Thr Asp 245 250 255
- Gly Asn Asp Thr Val Thr Glu Ser Ser Asp Gly Leu Trp Asn Asn Asn 260 265 270
- Gln Thr Gln Leu Phe Leu Glu His Ser Leu Leu Thr Ala Asn Thr Thr 275 280 285
- Lys Gly Ile Tyr Ala Gly Val Cys Ile Ser Val Leu Val Leu Leu Ala 290 295 300
- Leu Leu Gly Val Ile Ile Ala Lys Lys Tyr Phe Phe Lys Lys Glu Val 305 310 315 320
- Gln Gln Leu Arg Pro His Lys Ser Cys Ile His Gln Arg Glu 325 330

What is claimed is:

- 1. A purified and isolated DNA molecule having a nucleotide sequence set forth in SEQ ID
- 2 NO:1, SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6.
- 1 2. A purified and isolated DNA molecule selected from among:
- a) the DNA molecule of SEQ ID NO:1 or its complementary strand;
- b) the DNA molecule of SEO ID NO:2 or its complementary strand;
- 4 c) the DNA molecule of SEQ ID NO:4 or its complementary strand;
- 5 d) the DNA molecule of SEQ ID NO:6 or its complementary strand;
- 6 e) DNA molecules which hybridize under stringent conditions to the DNA molecule
- defined in a), b), c) or d), or fragments thereof;
- 8 f) DNA molecules which, but for the degeneracy of the genetic code, would hybridize to
- 9 the DNA molecule defined in a), b), c), d) or e).
- 3. The recombinant DNA molecule according to claim 1 or 2, operably linked to an
- 2 expression control sequence.
- 4. A vector comprising a purified and isolated DNA molecule having a nucleotide sequence
- 2 set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6.
- 5. A biologically functional plasmid or viral DNA vector comprising a DNA molecule
- 2 according to one of claims 1, 2 or 3.
- 6. A prokaryotic or eukaryotic host cell stably transformed or transfected by a vector
- 2 comprising a DNA molecule of claim 1.
- 7. A process for the production of a polypeptide product encoded by a DNA molecule
- 2 according to claim 1, 2 or 3, said process comprising:

- 49 -

- 3 growing, under suitable culture conditions, prokaryotic or eukaryotic host cells transformed
- 4 or transfected with the DNA molecule in a manner allowing expression of the DNA
- 5 molecule, and recovering the polypeptide product of said expression.
- 8. A polypeptide product produced by the process of claim 7.
- 9. A protein having an amino acid sequence which comprises SEQ ID NO:3, SEQ ID NO:5
- 2 or SEQ ID NO:7.
- 1 10. A purified and isolated protein encoded by the DNA of SEQ ID NO:1, SEQ ID NO:2,
- 2 SEQ ID NO:4 or SEQ ID NO:6.
- 1 11. The protein of claim 9 or 10, substantially free of other human proteins.
- 1 12. A protein which is a variant of SEQ ID NO:3, SEQ ID NO:5 or SEQ ID NO:7.
- 1 13. A soluble variant of the protein according to claim 9, 10, 11 or 12.
- 1 14. An IgG fusion protein comprising the protein of claim 9, 10, 11, 12 or 13.
- 1 15. The soluble protein of claim 13, fused to a toxin, imageable compound or radionuclide.
- 1 16. A specific monoclonal antibody to a protein of claim 9, 10, 11 or 12.
- 1 17. The antibody of claim 16, associated with a toxin, imageable compound or radionuclide.
- 1 18. A hybridoma cell line which produces a specific antibody to the protein of claim 9, 10,
- 2 11, 12 or 13.
- 1 19. An antibody produced by a hybridoma of claim 18.

- 20. A pharmaceutical composition comprising a therapeutically effective amount of the
- 2 protein of claim 9, 10, 11, 12, 13, 14 or 15, and further comprising a pharmacologically
- 3 acceptable carrier.

WO 97/44460

- 1 21. A pharmaceutical composition comprising a therapeutically effective amount of the
- 2 antibody of claim 16, 17 or 19, and further comprising a pharmacologically acceptable carrier.
- 22. A method of treating a subject with renal disease, comprising administering to the
- 2 subject a therapeutically effective amount of the protein of claim 9, 10, 11, 12, 13, 14 or 15.
- 1 23. A method of treating a subject with renal disease, comprising administering to the
- 2 subject a therapeutically effective amount of the antibody of claim 16, 17 or 19.
- 1 24. A method of treating a subject with renal disease, comprising administering to the
- 2 subject a therapeutically effective amount of the pharmaceutical composition of claim 20.
- 1 25. A method of promoting growth of new tissue in a subject, comprising administering to
- 2 the subject a therapeutically effective amount of the protein of claim 9, 10, 11, 12, 13 or 14.
- 1 26. The method of claim 25, wherein the tissue is renal tissue.
- 1 27. A method of promoting survival of damaged tissue in a subject, comprising
- 2 administering to the subject a therapeutically effective amount of the protein of claim 9, 10, 11,
- 3 12, 13 or 14.
- 1 28. The method of claim 27, wherein the tissue is renal tissue.
- 1 29. A method of treating a subject with renal disease, comprising administering to the
- 2 subject a therapeutically effective amount of the antibody of claim 16, 17 or 19.

- 51 -

- 30. A method of treating a subject with renal disease, comprising administering to the
 subject a therapeutically effective amount of the pharmaceutical composition of claim 21.
- 31. A method of promoting growth of new tissue in a subject, comprising administering to
 the subject a therapeutically effective amount of the antibody of claim 16, 17 or 19.
- 1 32. A method of promoting survival of damaged tissue in a subject, comprising
- 2 administering to the subject a therapeutically effective amount of the antibody of claim 16, 17 or
- 3 19.
- 1 33. A method of treating a subject with a renal disorder, comprising administering to the subject a vector of claim 4 or 5.
- 1 34. A method of promoting growth of new tissue in a subject, comprising administering to 2 the subject a vector of claim 4 or 5.
- 35. A method of promoting survival of damaged tissue in a subject, comprising
 administering a therapeutically effective amount of a vector of claim 4 or 5 to the subject.
- 1 36. The method of claim 34 or 35, wherein the tissue is renal tissue.
- 1 37. A method for targeting an imageable compound to a cell expressing a protein of SEO
- 2 ID NO:3, SEQ ID NO:5 or SEQ ID NO:7, comprising contacting the cell with a monoclonal
- 3 antibody of claim 16 fused to an imageable compound.
- 1 38. The method of claim 37, wherein the cell is within a subject, and the monoclonal
- 2 antibody is administered to the subject.
- 39. A method of identifying damage or regeneration of renal cells in a subject, comprising
- 2 comparing level of expression of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6

- 3 in renal cells of the subject to a control level of expression of SEQ ID NO:1, SEQ ID NO:2, SEQ
- 4 ID NO:4 or SEQ ID NO:6 in control renal cells.
- 40. A method of identifying upregulation of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:4
- 2 or SEQ ID NO:6 in cells comprising contacting the cells with an antisense probe and measuring
- 3 hybridization to RNA within the cell.
- 1 41. A method of identifying damage or regeneration of renal cells in a subject, comprising
- 2 comparing concentration of SEQ ID NO:3, SEQ ID NO:5 or SEQ ID NO:7 in renal cells, renal
- 3 cell fragments or body fluids of the subject to a control level of expression of SEQ ID NO:3,
- 4 SEQ ID NO:5 or SEQ ID NO:7 in control renal cells.
- 1 42. The method of claim 41, wherein the fluid is urine or serum.
- 1 43. The method of claim 41, wherein the renal cells or renal cell fragments are obtained
- 2 from urine sediment of the subject.

1	GCGGCCGCGTCGACGGTGCCTGTGAGTAAATAGATCAGGGTCTCCTTCAC	50
51	AGCACATTCTCCAGGAAGCCGAGCAAACATTAGTGCTATTTTACCCAGGA	100
101	GGAAATCTAGGTGTAGAGAGCTCTACGGATCTAAGGTTTGGATCTGTACC	150
151	CAGTGCTTTTTAGGTGTCTTTAGACATTTCTCAGGAAGATGTAGTCTCT	200
201	GTCACCATGTGTGGCTGAATTCTAGCTCAGTCCATCTTATTGTGTTTAAG	250
251	GTAGTTGAAGTTTAGGAACCAACCAGTATGTCTCTGAGCAGAAGAGTACA	300
301	GTGTCCATCTTGAGGACAAGCTCATCTTTACCATTAGAGGGCTGGCCTTG	350
351	GCTTAGATTCTACCGAGAACATACTCTCTAATGGCTGCCCTCAGTTTTCT	400
401	CTGTTTGCTGTCTTATTTGTGTCATGGCCAGAAGTCATATGGATGG	450
451	ATGTGAGCAAGGACCCAGATAGAAGAGTGTATTTGGGGGAACAGGTTGCC	500
501	CTAACAGAGAGTCCTGTGGGATTCATGCAGTCAGGATGAAGACCTGATCA	550
551	GACAGAGTGTGCTGAGTGCCACGGCTAACCAGAGTGACTTGTCACTGTCC	600
601	TTCAGGTCAACACCATGGTTCAACTTCAAGTCTTCATTTCAGGCCTCCTG M V Q L Q V F I S G L L	650
651	CTGCTTCTTCCAGGCTCTGTAGATTCTTATGAAGTAGTGAAGGGGGTGGT L L P G S V D S Y E V V K G V V	700
701	GGGTCACCCTGTCACAATTCCATGTACTTACTCAACACGTGGAGGAATCA G H P V T I P C T Y S T R G G I T	750
751	CAACGACATGTTGGGGCCGGGGCAATGCCCATATTCTAGTTGTCAAAAT T T C W G R G Q C P Y S S C Q N	800
801	ATACTTATTTGGACCAATGGATACCAAGTCACCTATCGGAGCAGCGGTCG I L I W T N G Y Q V T Y R S S G R	850
851	ATACAACATAAAGGGGCGTATTTCAGAAGGAGACGTATCCTTGACAATAG Y N I K G R I S E G D V S L T I E	900
901	AGAACTCTGTTGATAGTGATAGTGGTCTGTATTGTTGCCGAGTGGAGATT N S V D S D S G L Y C C R V E I	950
951	CCTGGATGGTTCAACGATCAGAAAATGACCTTTTCATTGGAAGTTAAACC P G W F N D Q K M T F S L E V K P	1000
1001	AGAAATTCCCACAAGTCCTCCAACAAGACCCACAACTACAAGACCCACAA E I P T S P P T R P T T R P T T	1050
1051	CCACAAGGCCCACAACTATTTCAACAAGATCCACACATGTACCAACATCA	1100

FIG. 1a

SUBSTITUTE SHEET (RULE 26)

1101 ACCAGAGTCTCCACCTCTACTCCAACACCAGAACAAACACAGACTCACAA 1150 TRVSTSTPTPEQTOTHK 1151 ACCAGAAATCACTACATTTTATGCCCATGAGACAACTGCTGAGGTGACAG 1200 PEITTFYAHETTAEVTE 1201 AAACTCCATCATATACTCCTGCAGACTGGAATGGCACTGTGACATCCTCA 1250 TPSYTPADWNGTVTSS GAGGAGGCCTGGAATAATCACACTGTAAGAATCCCTTTGAGGAAGCCGCA 1300 EEAWNNHTVRIPLRKPQ 1301 GAGAAACCCGACTAAGGGCTTCTATGTTGGCATGTCCGTTGCAGCCCTGC 1350 RNPTKGFYVGMSVAALL TGCTGCTGCTTGCGAGCACCGTGGTTGTCACCAGGTACATCATTATA 1351 1400 LLLLASTVVVTRYIII 1401 AGAAAGAAGATGGGCTCTCTGAGCTTTGTTGCCTTCCATGTCTCTAAGAG 1450 R K K M G S L S F V A F H V S K S TAGAGCTTTGCAGAACGCAGCGATTGTGCATCCCCGAGCTGAAGACAACA 1451 1500 R A L Q N A A I V H P R A E D N 1501 TCTACATTATTGAAGATAGATCTCGAGGTGCAGAATGAGTCCCAGAGGCC 1550 YIIEDRSRGAE TTCTGTGGGGCCTTCTGCCTGGGATTACAGAGATCGTGACTGATTTCACA 1551 1600 ${\tt GAGTAAAATACCCATTCCAGCTCCTGGGAGATTTTGTGTTTTTGGTTCTTC}$ 1601 1650 1651 CAGCTGCAGTGGAGAGGGTAACCCTCTACCCTGTATATGCAAAACTCGAG 1700 GTTAACATCATCCTAATTCTTGTATCAGCAACACCTCAGTGTCTCCACTC 1701 1750 ACTGCAGCGATTCTCTCAAATGTGAACATTTTAGAAGTTTGTGTTTCCTT 1751 1800 TTGTCCATGTAATCATTGGTAATACAAGAATTTTATCTTGTTTATTAAAA 1801 1850 1851 CCATTAATGAGAGGGGAATAGGAATTAAAAGCTGGTGGGAAGGGCCTCCT 1900 GAATTTAGAAGCACTTCATGATTGTGTTTATCTCTTTTTATTGTAATTTGA 1901 1950 AATGTTACTTCTATCCTTCCCAAGGGGCAAAATCATGGGAGCATGGAGGT 1951 2000 TTTAATTGCCCTCATAGATAAGTAGAAGAAGAGAGTCTAATGCCACCAAT 2050 AGAGGTGGTTATGCTTTCTCACAGCTCTGGAAATATGATCATTTATTATG 2051 2100 CAGTTGATCTTAGGATGAGGATGGGTTTCTTAGGAGGAGGGTTACCATG 2101 2150 GTGAGTGGACCAGGCACACATCAGGGGAAGAAAACAATGGATCAAGGGAT 2200 2201 TGAGTTCATTAGAGCCATTTCCACTCCACTTCTGTCTTGATGCTCAGTGT 2250 2251 TCCTAAACTCACCCACTGAGCTCTGAATTAGGTGCAGGGAGGAGACGTGC 2300

FIG. 1b

SUBSTITUTE SHEET (RULE 26)

3/11

2301	AGAAACGAAAGAGGAAAGGAGAGAGAGAGACACAGGCTTTCTGC	2350
2351	TGAGAGAAGTCCTATTGCAGGTGTGACAGTGTTTGGGACTACCACGGGTT	2400
2401	TCCTTCAGACTTCTAAGTTTCTAAATCACTATCATGTGATCATATTTATT	2450
2451	TTTAAAATTATTTCAGAAAGACACCACATTTTCAATAATAAATCAGTTTG	2500
2501	TCACAATTAATAAAATATTTTGTTTGCTAAGAAGTAAAAAAAA	2550
2551	AAGTCGACGCCGC 2566	

FIG. 1c

1	GCGGCCGCGTCGACGGTGCCTGTGAGTAAATAGATCAGGGTCTCCTTCAC	50
51	AGCACATTCTCCAGGAAGCCGAGCAAACATTAGTGCTATTTTACCCAGGA	100
101	GGAAATCTAGGTGTAGAGAGCTCTACGGATCTAAGGTCAACACCATGGTT M V	150
151	CAACTTCAAGTCTTCATTTCAGGCCTCCTGCTGCTTCTTCCAGGCTCTGT Q L Q V F I S G L L L L P G S V	200
201	AGATTCTTATGAAGTAGTGAAGGGGGTGGTGGGTCACCCTGTCACAATTC D S Y E V V K G V V G H P V T I P	250
251	CATGTACTTACTCAACACGTGGAGGAATCACAACGACATGTTGGGGCCGG C T Y S T R G G I T T T C W G R	300
301	GGGCAATGCCCATATTCTAGTTGTCAAAATATACTTATTTGGACCAATGG G Q C P Y S S C Q N I L I W T N G	350
351	ATACCAAGTCACCTATCGGAGCAGCGGTCGATACAACATAAAGGGGCGTA Y Q V T Y R S S G R Y N I K G R I	400
401	TTTCAGAAGGAGACGTATCCTTGACAATAGAGAACTCTGTTGATAGTGAT S E G D V S L T I E N S V D S D	450
451	AGTGGTCTGTATTGTTGCCGAGTGGAGATTCCTGGATGGTTCAACGATCA S G L Y C C R V E I P G W F N D Q	500
501	GAAAATGACCTTTTCATTGGAAGTTAAACCAGAAATTCCCACAAGTCCTC K M T F S L E V K P E I P T S P P	550
551	CAACAAGACCCACAACTACAAGACCCACAACCACAACCACTATT T R P T T T R P T T R P T T I	600
601	TCAACAAGATCCACACATGTACCAACATCAACCAGAGTCTCCACCTCTAC S T R S T H V P T S T R V S T S T	650
651	TCCAACACCAGAACAAACAGACTCACAAACCAGAAATCACTACATTTT P T P E Q T Q T H K P E I T T F Y	700
701	ATGCCCATGAGACAACTGCTGAGGTGACAGAAACTCCATCATATACTCCT A H E T T A E V T E T P S Y T P	750
751	GCAGACTGGAATGGCACTGTGACATCCTCAGAGGAGGCCTGGAATAATCA A D W N G T V T S S E E A W N N H	800
801	CACTGTAAGAATCCCTTTGAGGAAGCCGCAGAGAAACCCGACTAAGGGCT T V R I P L R K P Q R N P T K G F	850
851	TCTATGTTGGCATGTCCGTTGCAGCCCTGCTGCTGCTGCTGCTGCGAGC Y V G M S V A A L L L L L A S	900
901	ACCGTGGTTGTCACCAGGTACATCATTATAAGAAAGAAGATGGGCTCTCT T V V V T R Y I I I R K K M G S L	950

FIG. 2a SUBSTITUTE SHEET (RULE 26)

951	GAGCTTTGTTGCCTTCCATGTCTCTAAGAGTAGAGCTTTGCAGAACGCAG S F V A F H V S K S R A L Q N A A	1000
1001	CGATTGTGCATCCCCGAGCTGAAGACAACATCTACATTATTGAAGATAGA I V H P R A E D N I Y I I E D R	1050
1051	TCTCGAGGTGCAGAATGAGTCCCAGAGGCCTTCTGTGGGGCCTTCTGCCT S R G A E	1100
1101	GGGATTACAGAGATCGTGACTGATTTCACAGAGTAAAATACCCATTCCAG	1150
1151	CTCCTGGGAGATTTTGTGTTTTTGGTTCTTCCAGCTGCAGTGGAGAGGGTA	1200
1201	ACCCTCTACCCTGTATATGCAAAACTCGAGGTTAACATCATCCTAATTCT	1250
1251	TGTATCAGCAACACCTCAGTGTCTCCACTCACTGCAGCGATTCTCTCAAA	1300
1301	TGTGAACATTTTAGAAGTTTGTGTTTTCCTTTTGTCCATGTAATCATTGGT	1350
1351	AATACAAGAATTTTATCTTGTTTATTAAAACCATTAATGAGAGGGGAATA	1400
1401	GGAATTAAAAGCTGGTGGGAAGGGCCTCCTGAATTTAGAAGCACTTCATG	· 1450
1451	ATTGTGTTTATCTCTTTTATTGTAATTTGAAATGTTACTTCTATCCTTCC	1500
1501	CAAGGGGCAAAATCATGGGAGCATGGAGGTTTTAATTGCCCTCATAGATA	1550
1551	AGTAGAAGAAGAGAGTCTAATGCCACCAATAGAGGTGGTTATGCTTTCTC	1600
1601	ACAGCTCTGGAAATATGATCATTTATTATGCAGTTGATCTTAGGATGAGG	1650
1651	ATGGGTTTCTTAGGAGGAGGGTTACCATGGTGAGTGGACCAGGCACACA	1700
1701	TCAGGGGAAGAAAACAATGGATCAAGGGATTGAGTTCATTAGAGCCATTT	1750
1751	CCACTCCACTTCTGTCTTGATGCTCAGTGTTCCTAAACTCACCCACTGAG	1800
1801	CTCTGAATTAGGTGCAGGAGGGAGAGAGAGAGAGAGAGAG	1850
1851	AGGAGAGAGAGACACAGGCTTTCTGCTGAGAGAAGTCCTATTGCAG	1900
1901	GTGTGACAGTGTTTGGGACTACCACGGGTTTCCTTCAGACTTCTAAGTTT	1950
1951	CTAAATCACTATCATGTGATCATATTTATTTTTAAAATTATTTCAGAAAG	2000
2001	ACACCACATTTTCAATAATAAATCAGTTTGTCACAATTAATAAAATATTT	2050
2051	TGTTTGCTAAGAAGTAAAAAGTCGACGCGGCCGC 2084	

FIG. 2b

1	GCGGCCGCGTCGACTCGCAGGAGGCCGGCACTCTGACTCCTGGTGGATGG	50
51	GACTAGGGAGTCAGAGTCAAGCCCTGACTGGCTGAGGGCGGCGCTCCGA	100
101	GTCAGCATGGAAAGTCTCTGCGGGGTCCTGGTATTTCTGCTGCTGCTGC M E S L C G V L V F L L L A A	150
151	AGGACTGCCGCTCCAGGCGGCCAAGCGGTTCCGTGATGTGCTGGGCCATG G L P L Q A A K R F R D V L G H E	.200
201	AGCAGTATCCGGATCACATGAGGGAGAACAACCAATTACGTGGCTGGTCT Q Y P D H M R E N N Q L R G W S	250
251	TCAGATGAAAATGAATGGGATGAACAGCTGTATCCAGTGTGGAGGAGGGG S D E N E W D E Q L Y P V W R R G	300
301	AGAGGGCAGATGGAAGGACTCCTGGGAAGGAGGCCGTGTGCAGGCAG	350
351	TAACCAGTGATTCACCGGCCTTGGTGGGTTCCAATATCACCTTCGTAGTG T S D S P A L V G S N I T F V V	400
401	AACCTGGTGTTCCCCAGATGCCAGAGGAAGATGCCAACGGCAATATCGT N L V F P R C Q K E D A N G N I V	450
451	CTATGAGAGGAACTGCAGAAGTGATTTGGAGCTGGCTTCTGACCCGTATG Y E R N C R S D L E L A S D P Y V	500
501	TCTACAACTGGACCACAGGGCAGACGACGACGCACC Y N W T T G A D D E D W E D S T	550
551	AGCCAAGGCCAGCACCTCAGGTTCCCCGACGGGAAGCCCTTCCCTCGCCC S Q G Q H L R F P D G K P F P R P	600
601	CCACGGACGAAGAAATGGAACTTCGTCTACGTCTTCCACACACTTGGTC H G R K K W N F V Y V F H T L G Q	650
651	AGTATTTTCAAAAGCTGGGTCGGTGTTCAGCACGAGTTTCTATAAACACA Y F Q K L G R C S A R V S I N T	700
701	GTCAACTTGACAGTTGGCCCTCAGGTCATGGAAGTGATTGTCTTTCGAAG V N L T V G P Q V M E V I V F R R	750
751	ACACGGCCGGCATACATTCCCATCTCCAAAGTGAAAGACGTGTATGTGA H G R A Y I P I S K V K D V Y V I	800
801	TAACAGATCAGATCCCTATATTCGTGACCATGTACCAGAAGAATGACCGG T D Q I P I F V T M Y Q K N D R	850
851	AACTCGTCTGATGAAACCTTCCTCAGAGACCTCCCCATTTTCTTCGATGT N S S D E T F L R D L P I F F D V	900
901	CCTCATTCACGATCCCAGTCATTTCCTCAACTACTCTGCCATTTCCTACA	950

FIG. 3a

SUBSTITUTE SHEET (RULE 26)

951	AGTGGAACTTTGGGGACAACACTGGCCTGTTTGTCTCCAACAATCACACTW N F G D N T G L F V S N N H T	1000
1001	TTGAATCACACGTATGTGCTCAATGGAACCTTCAACTTTAACCTCACCGT L N H T Y V L N G T F N F N L T V	1050
1051	GCAAACTGCAGTGCCGGGACCATGCCCCTCACCCACACCTTCGCCTTCTT Q T A V P G P C P S P T P S P S S	1100
1101	CTTCGACTTCTCCTTCGCCTGCATCTTCGCCTTCACCCACATTATCAACA S T S P S P A S S P S P T L S T	1150
1151	CCTAGTCCCTCTTTAATGCCTACTGGCCACAAATCCATGGAGCTGAGTGA PSPSLMPTGHKSMELSD	1200
1201	CATTTCCAATGAAAACTGCCGAATAAACAGATATGGTTACTTCAGAGCCA I S N E N C R I N R Y G Y F R A T	1250
1251	CCATCACAATTGTAGATGGAATCCTAGAAGTCAACATCATCCAGGTAGCA I T I V D G I L E V N I I Q V A	1300
1301	GATGTCCCAATCCCCACACCGCAGCCTGACAACTCACTGATGGACTTCAT D V P I P T P Q P D N S L M D F I	1350
1351	TGTGACCTGCAAAGGGGCCACTCCCACGGAAGCCTGTACGATCATCTCTG V T C K G A T P T E A C T I I S D	1400
1401	ACCCCACCTGCCAGATCGCCCAGAACAGGGTGTGCAGCCCGGTGGCTGTG P T C Q I A Q N R V C S P V A V	1450
1451	GATGAGCTGTGCCTCCTGTCCGTGAGGAGCCTTCAATGGGTCCGGCAC D E L C L L S V R R A F N G S G T	1500
1501	GTACTGTGTGAATTTCACTCTGGGAGACGATGCAAGCCTGGCCCTCACCA Y C V N F T L G D D A S L A L T S	1550
1551	GCGCCTGATCTCTATCCCTGGCAAAGACCTAGGCTCCCCTCTGAGAACA A L I S I P G K D L G S P L R T	1600
1601	GTGAATGGTGTCCTGATCTCCATTGGCTGCCTGGCCATGTTTGTCACCAT V N G V L I S I G C L A M F V T M	1650
1651	GGTTACCATCTTGCTGTACAAAAAACACAAGACGTACAAGCCAATAGGAA V T I L L Y K K H K T Y K P I G N	1700
1701	ACTGCACCAGGAACGTGGTCAAGGGCAAAGGCCTGAGTGTTTTTCTCAGC C T R N V V K G K G L S V F L S	1750
1751	CATGCAAAAGCCCCGTTCTCCCGAGGAGACCGGGAGAAGGATCCACTGCT H A K A P F S R G D R E K D P L L	1800
1801	CCAGGACAAGCCATGGATGCTCTAAGTCTTCACTCTCACTTCTGACTGGG Q D K P W M L	1850
1851	AACCCACTCTTCTCTCCATCTATCTCACCTCTCCACAACTACATCACTCC	1000

FIG. 3b

SUBSTITUTE SHEET (RULE 26)

1901	TAGCTGTTGTTTTCTACGGATTATTGTAAAATGTATATCATGGTTTAGGG	1950
1951	AGCGTAGTTAATTGGCATTTTAGTGAAGGGATGGGAAGACAGTATTTCTT	2000
2001	CACATCTGTATTGTGGTTTTATACTGTTAATAGGGTGGGCACATTGTGT	2050
2051	CTGAAGGGGGAGGGGAGGTCACTGCTACTTAAGGTCCTAGGTTAACTGG	2100
2101	GAGAGGATGCCCCAGGCTCCTTAGATTTCTACACAAGATGTGCCTGAACC	2150
2151	CAGCTAGTCCTGACCTAAAGGCCATGCTTCATCAACTCTATCTCAGCTCA	2200
2201	TTGAACATACCTGAGCACCTGATGGAATTATAATGGAACCAAGCTTGTTG	2250
2251	TATGGTGTGTGTGTACATAAGATACTCATTAAAAAGACAGTCTATTAA	2300
2301	AAA 2303	

FIG. 3c

1	ATGCATCCTCAAGTGGTCATCTTAAGCCTCATCCTACATCTGGCAGATTC	50
	M H P Q V V I L S L I L H L A D S	
51	TGTAGCTGGTTCTGTAAAGGTTGGTGGAGAGGCAGGTCCATCTGTCACAC V A G S V K V G G E A G P S V T L	100
101	TACCCTGCCACTACAGTGGAGCTGTCACATCAATGTGCTGGAATAGAGGC P C H Y S G A V T S M C W N R G	150
151	TCATGTTCTCTATTCACATGCCAAAATGGCATTGTCTGGACCAATGGAAC S C S L F T C Q N G I V W T N G T	200
201	CCACGTCACCTATCGGAAGGACACACGCTATAAGCTATTGGGGGACCTTT H V T Y R K D T R Y K L L G D L S	250
251	CAAGAAGGGATGTCTCTTTGACCATAGAAAATACAGCTGTGTCTGACAGT R R D V S L T I E N T A V S D S	300
301	GGCGTATATTGTTGCCGTGTTGAGCACCGTGGGTGGTTCAATGACATGAA G V Y C C R V E H R G W F N D M K	350
351	AATCACCGTATCATTGGAGATTGTGCCACCCAAGGTCACGACTACTCCAA I T V S L E I V P P K V T T T P I	400
401	TTGTCACAACTGTTCCAACCGTCACGACTGTTCGAACGAGCACCACTGTTVTTVTTVRTSTTV	450
451	CCAACGACAACGACTGTTCCAACGACAACAACAATGAGCAT P T T T V P T T V P T T M S I	500
501	TCCAACGACAACGACTGTTCCGACGACAATGACTGTTTCAACGACAACGA P T T T V P T T M T V S T T T S	550
551	GCGTTCCAACGACAACGAGCATTCCAACAACAACAAGTGTTCCAGTGACA V P T T T S I P T T T S V P V T	600
601	ACAACGGTCTCTACCTTTGTTCCTCCAATGCCTTTGCCCAGGCAGAACCA T T V S T F V P P M P L P R Q N H	650
651	TGAACCAGTAGCCACTTCACCATCTTCACCTCAGCCAGCAGAAACCCACC E P V A T S P S S P Q P A E T H P	700
701	CTACGACACTGCAGGGAGCAATAAGGAGAACCCACCAGCTCACCATTG T T L Q G A I R R E P T S S P L	750
751	TACTCTTACACAACAGATGGGAATGACACCGTGACAGAGTCTTCAGATGG Y S Y T T D G N D T V T E S S D G	800
801	CCTTTGGAATAACAATCAAACTCAACTGTTCCTAGAACATAGTCTACTGA L W N N N Q T Q L F L E H S L L T	850
851	CGGCCAATACCACTAAAGGAATCTATGCTGGAGTCTGTATTTCTGTCTTG	900

FIG. 4a

901	GTGCTTCTTGCTCTTTTGGGTGTCATCATTGCCAAAAAGTATTTCTTCAA												950					
	V	L	L	A	\mathbf{L}	L	G	V	1	I	A	K	K	Y	F	F	K	
051	AAA		እሶሶ	ጥጥ⁄	7 7 C	א א <i>א</i>	תמיד	~ h ~ /	2001	A TTO TA	• • • • • • • • • • • • • • • • • • • •		ת יחיי		a mo			1000
321																	GAG E	1000
1001	AA	10	002	_	_												_	

FIG. 4b

1	MHPQVVILSLILHLADSVAGSVKVGGEAGPSVTLPCHYSGAVTSMCWN	48
2	:: . : : :: .: 	51
49	RGSCSLFTCONGIVWTNGTHVTYRKDTRYKLLGDLSRRDVSLTIENTAVS	98
52	RGQCPYSSCQNILIWTNGYQVTYRSSGRYNIKGRISEGDVSLTIENSVDS	101
99	DSGVYCCRVEHRGWFNDMKITVSLEIVPPKVTTTPIVTTVPTVTTVRTST	148
102	: . : . :	135
149	TVPTTTVPTTMSIPTTTVPTTMTVSTTTSVPTTTSIPTTTSVP	198
136	PTRPTTTRPTTTISTRSTHVPTSTRVSTSTPTPEQTQTHKP	180
199	VTTTVSTFVPPMPLPRQNHEPVATSPSSPQPAETHPTTLQGAIRREPTSS	248
181	eittfyahettaevtetp	198
249	PLYSYTTDGNDTVTESSDGLWNNNQTQLFLEHSLLTANTTKGIYAGVCIS	298
199	:: :: :: . : ::::	243
299	VLVLLALLGVIIAKKY.FFKKEVQQLRPHKSCIHQRE 3	34
244	ALLLLLASTVVVTRYIIIRKKMGSLSFVAFHVSKSRALONAAIVHPRA 2	92

FIG. 5

INTERNATIONAL SEARCH REPORT

Internatic. Application No PCT/US 97/89383

			<u> </u>						
A CLASSE IPC 6	FICATION OF SUBJECT MATTER C12N15/12 C07K14/47 C12N15/ G01N33/50 C12Q1/68 C12N1/2 A61K48/00	62 C07K16/18 1 C12N5/10	A61K38/16 C12N5/12						
According to	International Patent Classification (IPC) or to both national classific	ation and IPC							
	SEARCHED								
Minimum do	cumentation searched (classification system followed by classification	on symbols)							
IPC 6	C12N C07K A61K G01N C12Q								
Documental	Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched								
	ata base consulted during the international search (name of data ba	se and, where practical, seen	ch terms used)						
	ENTS CONSIDERED TO BE RELEVANT								
Category *	Citation of document, with indication, where appropriate, of the rel	event pessages	Relevant to claim No.						
х	WO 96 04376 A (US HEALTH) 15 Feb see page 39 - page 41 see page 43	ruary 1996	1						
X	WETERMAN A.M.J. ET AL: "nmb, a gene, is expressed in low-metast melanoma cell lines and xenograf INT. J. CANCER, vol. 60, 1995, pages 73-81, XP002044250 see the whole document	atic human	1						
Furth	ner documents are listed in the continuation of box C.	X Patent family memb	ers are listed in annex.						
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A docume	nt defining the general state of the art which is not	or priority data and not i	n conflict with the application but principle or theory underlying the						
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22	2 October 1997	1	4. 11. 97						
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	European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl.								
	Fax: (+31-70) 340-3018	Espen, J							

PCT/US 97/09303

INTERNATIONAL SEARCH REPORT

The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.	amoA
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Claims Mos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a). Observations where unity of invention is laciding (Continuation of item 2 of first sheet)	3. E
Cleims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:	
Claims Nos.: See FURTHER INFORMATION sheet pe searched by this Authority, namely: See FURTHER INFORMATION sheet PCT/ISA/210	X .
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INTERNATIONAL SEARCH REPORT

International Application No. PCT/IS 97 09303

FURTHER INFORMATION CONTINUED FROM PO	:T/ISA/	21A
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HTHER INFORMATION CONTINUED FROM PCT/ISA/ 210	
Remark: Although claims 22-36 are directed to a method of treatment of the human/animal body, and although claims 38-39, and in part 37,40,41 are directed to a diagnostic method practised on the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.	
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INTERNATIONAL SEARCH REPORT

International Application No PCT/US 97/09303

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Patent document cited in search report	Publication date	Patent far member			Publication date
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